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(57) Abstract

Activation by CD40 ligand of renal cells bearing CD40 on the cell surface is inhibited, both in vivo and ex vivo, with an agent capable of inhibiting interaction between CD40 ligand and CD40 on the cells. Inflammatory kidney diseases are treated.

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THERAPEUTIC APPLICATIONS OF T-BAM (CD40L) TECHNOLOGY TO TREAT INFLAMMATORY KIDNEY DISEASES

This application claims the priority of U.S. Serial No. 08/641,473, filed May 1, 1996 and U.S. Serial No. 08/587,334, filed January 16, 1996, the contents of which are hereby incorporated by reference.

The invention disclosed herein was made with Government support under NIH Grant Nos. K08-AR-01904, R01-CA55713, R01-AI-28367, R01-AI-14969, HL21006, HL42833, HL50629, and R01-AI-14969 from the Department of Health and Human Services. Accordingly, the U.S. Government has certain rights in this invention.

Throughout this application, various references are referred to within parentheses. Disclosures of these publications in their entireties are hereby incorporated by reference into this application to more fully describe the state of the art to which this invention pertains. Full bibliographic citation for these references may be found in the text.

Background of the Invention

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Immune complex deposition is known to play important roles in mediating the immunopathogenesis of a variety of glomerulonephritis renal diseases, including the associated with systemic lupus erythematosus. infiltrating renal interstitial leukocytes, predominately T cells and monocytes, are often seen in lupus nephritis and other inflammatory renal diseases. The precise role of infiltrating T cells in the inflammatory renal process ultimately may result in renal scarring and endorgan damage is currently unknown. It is of interest that the extent of mononuclear cell infiltrate correlates with progression to renal failure. Some evidence

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suggests that interstitial T cells play direct immunopathogenic roles in the initiation and/or propagation of inflammatory renal diseases, including lupus nephritis.

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CD40 is a cell surface molecule expressed on a variety of cells and interacts with a 30-33 kDa activation-induced CD4+ T cell counterreceptor termed CD40L. CD40L-CD40 interactions have been extensively studied in T cell-B cell interactions and are essential for T cell dependent B cell differentiation and IgG, IgA and IgE production. CD40 is also expressed on monocytes, dendritic cells, epithelial cells, endothelial cells and fibroblasts. CD40 expression on these cells is upregulated in vitro by cytokines, most notably IFN-y. In vivo studies have demonstrated markedly upregulated CD40 expression in inflammatory sites, such as rheumatoid arthritis synovial membrane or psoriatic plaques. In vitro studies utilizing anti-CD40 mAb or CD40L+ cells demonstrate that CD40 is functionally expressed on monocytes, dendritic cells, epithelial cells, endothelial cells and fibroblasts.

Earlier disclosure of treating idiopathic autoimmune 25 diseases, including drug-induced lupus, such as Patent Publication International No. WO 93/09812 (published May 27, 1993) was based on the finding that CD40 is expressed on the surface of B cells. initiation point of lupus is the deposition 30 autoantibodies in the kidney, which then attracts cells involved in destruction of kidney tissue. The finding, discussed below, that CD40 is expressed on kidney tubule cells provides the basis for treating inflammatory kidney diseases having initiation points other than autoantibody 35 deposition.

Summary of the Invention

This invention provides a method of inhibiting activation by CD40 ligand of renal cells bearing CD40 on the surface of the cells, comprising contacting the cells with an agent capable of inhibiting interaction between CD40 ligand and CD40 on the cells, the agent being present in an amount effective to inhibit activation of the cells.

This invention provides a method of inhibiting activation by CD40 ligand of renal cells bearing CD40 on the surface of the cells, in a subject, comprising administering to the subject an agent capable of inhibiting interaction between CD40 ligand and CD40 on the cells, the agent being present in an amount effective to inhibit activation of the cells in the subject.

This invention provides a method of treating, in a subject, an inflammatory kidney disease, comprising administering to the subject an agent capable of inhibiting interaction between CD40 ligand and CD40 on the cells, the agent being present in an amount effective to inhibit activation of the cells in the subject and thereby treat the inflammatory kidney disease.

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Description of the Figures

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Figures 1A-Y: Atomic coordinates of crystal structure of soluble extracellular fragment of human CD40L containing residues Gly116-Leu261 (in Brookhaven Protein Data Bank format). (SEQ ID NO:1).

Figures 2A-C: Expression of CD40 in normal kidney. 10 Shown are frozen sections of normal kidney stained with control mouse IgG (Figure 2A, magnification 25x) or anti-CD40 mAb G28.5 (Figures 2B and 2C, magnification 40x). Distal tubules and interstitial 15 capillaries express CD40 while proximal tubules are CD40 (Figure 2B). Glomerular cells and epithelial cells of Bowmans capsule express CD40 (Figure 2C).

20 Pigures 3A-C: Expression of CD40 in diffuse proliferative lupus nephritis. Shown are frozen sections of a kidney biopsy from a patient with Class IV lupus nephritis stained with control mouse IgG (Figure 3A, 25 magnification 25x) or anti-CD40 mAb G28.5 (Figures 3B and 3C, magnification 40x). Figure 3B shows intense CD40 staining of distal and proximal tubules. Figure 3C shows increased and diffuse CD40 30 expression in the glomerulus. Figure 3C also shows that the epithelial derived crescent is CD40+.

Figure 4A: CD40L expression on interstitial mononuclear cells in class IV lupus glomerulonephritis. Shown is a frozen section obtained from a renal biopsy

specimen stained with anti-CD40L mAb 5c8. Bound antibody was visualized with the Vectastain ABC Elit kit followed by the chromogen 3-amino-9-ethylcarbazole (Vector Laboratories). The tissue was counterstained with Mayer's hematoxylin (Sigma). CD40L immunoreactivity is noted as staining of mononuclear cells.

10 Figure 4B:

Isotype control staining of interstitial mononuclear cells in class IV lupus glomerulonephritis. Shown is a frozen section obtained from the same patient studied in Figure 4A and stained with an IgG2a isotype control mAb. Bound antibody was visualized with the Vectastain ABC Elite kit followed by the chromogen 3amino-9-ethylcarbazole (Vector tissue was The Laboratories). counterstained with Mayer's hematoxylin (Sigma). Note the lack of immunoreactivity (staining).

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Figure 5:

interstitial on CD40L expression mononuclear cells in class Shown is a frozen glomerulonephritis. section obtained from a renal biopsy specimen stained with anti-CD40L mAb 5c8. obtained was This specimen different patient than shown in Figure 4A. Bound antibody was visualized with the Vectastain ABC Elite kit followed by the chromogen 3-amino-9-ethylcarbazole (Vector tissue Laboratories). The counterstained with Mayer's hematoxylin (Sigma). CD40L immunoreactivity is noted cells. mononuclear staining of

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Staining with an isotype control mAb was negative (not shown).

Figure 6:

Renal CD40 expression in focal segmental glomerulosclerosis (FSGS). Shown is a frozen section obtained from a renal biopsy specimen stained with anti-CD40 mAb G28.5. Bound antibody was visualized with the Vectastain ABC Elite kit followed by the chromogen 3-amino-9-ethylcarbazole (Vector Laboratories). The tissue was counterstained with Mayer's hematoxylin (Sigma). Note the intense CD40 staining. Staining with an isotype control mAb was negative (not shown).

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Figure 7:

CD40L expression on interstitial in focal mononuclear cells segmental glomerulosclerosis. Shown is a frozen section obtained from the same patient as studied in Figure 6 stained with anti-5c8. CD40L mAb Bound antibody was visualized with the Vectastain ABC Elite kit followed by the chromogen 3-amino-9ethylcarbazole (Vector Laboratories). The tissue was counterstained with Mayer's hematoxylin (Sigma). immunoreactivity is noted as staining of mononuclear cells. Staining with an isotype control mAb was negative (not shown).

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Figure 8:

Renal CD40 expression in IgA nephropathy. Shown is a frozen section obtained from a renal biopsy specimen stained with anti-CD40 mAb G28.5. Bound antibody was visualized with the Vectastain ABC Elite

kit followed by the chromogen 3-amino-9-ethylcarbazole (Vector Laboratories). The tissue was counterstained with Mayer's hematoxylin (Sigma). Note the intense CD40 staining. Staining with an isotype control mAb was negative (not shown).

Figure 9:

interstitial expression on CD40L mononuclear cells in IgA nephropathy. Shown is a frozen section obtained from the same patient as studied in Figure 8 stained with anti-CD40L mAb 5c8. the visualized with antibody was Vectastain ABC Elite kit followed by the chromogen 3-amino-9-ethylcarbazole (Vector The tissue Laboratories). counterstained with Mayer's hematoxylin (Sigma). CD40L immunoreactivity is noted cells. of mononuclear as staining Staining with as isotype control mAb was negative (not shown).

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Detail d Description

This invention provides a method of inhibiting activation by CD40 ligand of renal cells bearing CD40 on the cell surface, comprising contacting the cells with an agent capable of inhibiting interaction between CD40 ligand and CD40 on the cells, the agent being present in an amount effective to inhibit activation of the cells. embodiment of this invention the agent is capable of inhibiting any interaction between CD40 ligand and CD40. "Interaction between CD40 ligand and CD40 on the cells" refers to one or more aspects, functional or structural, of a CD40-CD40 ligand interrelationship. Therefore, in one embodiment, an agent which inhibits interaction may competitively bind to CD40 ligand in such a way to block or diminish the binding of CD40 ligand to cellular CD40. In another embodiment an agent which inhibits interaction may associate with CD40 or CD40 ligand in a manner which does not inhibit binding of CD40 ligand to cellular CD40, but which influences the cellular response to the CD40 ligation, such as by altering the turnover rate of the cellular CD40 or the CD40-agent complex, by altering binding kinetics of CD40 with CD40 ligand, or by altering the rate or extent of cellular activation in response to CD40 ligation.

In specific embodiments the CD40-bearing renal cells are selected from the group consisting of glomerular endothelial cells, mesangial cells, distal tubule cells, proximal tubule cells, parietal epithelial cells, visceral epithelial cells, cells of a Henle loop or limb thereof, and interstitial inflammatory cells. In a more specific embodiment the parietal epithelial cells are crescent parietal epithelial cells.

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In an embodiment of this invention the agent inhibits binding of CD40 ligand to CD40 on the cells.

In an embodiment of this invention the agent is a protein.

In another embodiment of this invention the agent is a nonprotein. As used herein the term nonprotein includes 5 any and all compounds or agents which encompass elements other than simple or conjugated polypeptide chains. includes elements such as amino acids having non-peptide linkages; nonprotein amino acids such as β , γ , or δ amino acids, amino acids in D configuration, or other 10 acids homocysteine, including nonprotein amino homoserine, citrulline, ornithine, y-aminobutyric acid, β -cyanoalanine; djenkolic acid, or canavanine, carbohydrate polysaccharides, or monosaccharides, moieties; fatty acids or lipid moieties; nucleotide 15 moieties, mineral moieties; or other nonprotein elements.

In a specific embodiment the protein comprises an antibody or portion thereof capable of inhibiting 20 interaction between CD40 ligand and CD40 on the cells. The antibody is a monoclonal or polyclonal antibody. a more specific embodiment the monoclonal antibody specifically binds to the epitope to which monoclonal antibody 5c8 (ATCC Accession No. HB 10916) specifically 25 An example of such a monoclonal antibody is monoclonal antibody 5c8 (ATCC Accession No. HB 10916). In another embodiment, the antibody specifically binds to One example of an anti-CD40 antibody is the CD40. monoclonal mouse anti-human CD40, available from Genzyme 30 Customer Service (Product 80-3702-01, Cambridge, MA). other embodiments the monoclonal antibody is a chimeric antibody, a primatized antibody, a humanized antibody, or an antibody which includes a CDR region from a first human and an antibody scaffold from a second human. 35

The meaning of "chimeric", "primatized" and "humanized"

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antibody and methods of producing them are well known to those of skill in the art. See, for example, PCT International Publication No. WO 90/07861, published July 26, 1990 (Queen, et al.); and Queen, et al. Proc. Nat'l Acad. Sci.-USA (1989) 86: 10029). Methods of making primatized antibodies are disclosed, for example, in PCT International publication No. WO/02108, corresponding to International Application No. PCT/US92/06194 (Idec Pharmaceuticals); and in Newman, et al., Biotechnology (1992) 10:1455-1460, which are hereby incorporated by reference into this application.

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Generally, a humanized antibody is an antibody comprising one or more complementarity determining regions (CDRs) of antibody functionally joined to human a non-human framework region segments. Additional residues associated with the non-human antibody can optionally be Typically, at least one heavy chain or one present. light chain comprises non-human CDRs. Typically, the non-human CDRs are mouse CDRs. Generally, a primatized antibody comprising one antibody is an complementarity determining regions (CDRs) of an antibody of a species other than a non-human primate, functionally joined to framework region segments of a non-human primate. Additional residues associated with the species from which the CDR is derived can optionally be present. Typically, at least one heavy chain or one light chain comprises CDRs of the species which is not a nonhuman Typically, the CDRs are human CDRs. Generally, a chimeric antibody is an antibody whose light and/or heavy chains contain regions from different species. example one or more variable (V) region segments of one species may be joined to one or more constant (C) region segments of another species. Typically, a chimeric antibody contains variable region segments of a mouse joined to human constant region segments, although other mammalian species may be used.

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Monoclonal antibody 5c8 is produced by a hybridoma cell which was deposited on November 14, 1991 with the American Type Culture Collection (ATCC), 12301 Parklawn Drive, Rockville, Maryland 20852, U.S.A. under the provisions of the Budapest Treaty for the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure. The hybridoma was accorded ATCC Accession Number HB 10916.

In a specific embodiment the portion of the antibody comprises a complementarity determining region or variable region of a light or heavy chain. In another specific embodiment the portion of the antibody comprises a complementarity determining region or a variable region. In another specific embodiment the portion of the antibody comprises a Fab or a single chain antibody. A single chain antibody is made up of variable regions linked by protein spacers in a single protein chain.

In another embodiment the protein comprises soluble extracellular region of CD40 ligand, or portion thereof, or variant thereof, capable of inhibiting any interaction between CD40 ligand and CD40 on the cells; or soluble extracellular region of CD40, or portion thereof, or variant thereof, capable of inhibiting any interaction between CD40 ligand and CD40 on the cells. In a specific embodiment the soluble extracellular region of CD40 ligand or CD40 is a monomer. In another embodiment the soluble extracellular region of CD40 is an oligomer.

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Variants can differ from naturally occurring CD40 or CD40 ligand in amino acid sequence or in ways that do not involve sequence, or both. Variants in amino acid sequence are produced when one or more amino acids in naturally occurring CD40 or CD40 ligand is substituted with a different natural amino acid, an amino acid derivative or non-native amino acid. Particularly

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preferred variants include naturally occurring CD40 or ligand, or biologically active fragments naturally occurring CD40 or CD40 ligand, whose sequences differ from the wild type sequence by one or more conservative amino acid substitutions, which typically have minimal influence on the secondary structure and hydrophobic nature of the protein or peptide. may also have sequences which differ by one or more nonconservative amino acid substitutions, deletions or insertions which do not abolish the CD40 or CD40 ligand biological activity. Conservative substitutions typically include the substitution of one amino acid for another with similar characteristics such as substitutions within the following groups: valine, glycine; glycine, alanine; valine, isoleucine; aspartic acid, glutamic asparagine, glutamine; serine, threonine; lysine, arginine; and phenylalanine, tyrosine. The non-polar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine. polar neutral amino acids include glycine, threonine, cysteine, tyrosine, asparagine and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic) amino acids include aspartic acid and glutamic acid.

Other conservative substitutions can be taken from Table 1, and yet others are described by Dayhoff in the Atlas of Protein Sequence and Structure (1988).

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Table 1: Conservative Amino Acid Replacements

For Amino Acid	Code	Replace with any of
Alanine	A	D-Ala, Gly,beta-ALa, L-Cys,D- Cys
Arginine	R	D-Arg, Lys, homo-Arg, D-homo-Arg, Met, D-Met, Ile, D-Ile, Orn, D-Orn
Asparagine	N	D-Asn, Asp, D-Asp, Glu, D-Glu, Gln, D-Gln
Aspartic Acid	D	D-Asp,D-Asn,Asn, Glu,D-Glu, Gln, D-Gln
Cysteine	С	D-Cys, S-Me-Cys, Met, D-Met, Thr, D-Thr
Glutamine	Q	D-Gln, Asn, D-Asn, Glu, D-Glu, Asp, D-Asp
Glutamic Acid	E	D-Glu, D-Asp, Asp, Asn, D-Asn, Gln, D-Gln
Glycine	G	Ala, D-Ala, Pro, D-Pro, Beta- Ala, Acp
Isoleucine	I	D-Ile, Val, D-Val, Leu, D-Leu, Met, D-Met
Leucine	L	D-Leu, Val, D-Val, Met, D-Met
Lysine	K	D-Lys, Arg, D-Arg, homo-Arg, D-homo-Arg, Met, D-Met, Ile, D-Ile, Orn, D-Orn
Methionine	M	D-Met, S-Me-Cys, Ile, D-Ile, Leu, D-Leu, Val, D-Val, Norleu
Phenylalanine	F	D-Phe, Tyr, D-Thr, L-Dopa, His, D-His, Trp, D-Trp, Trans 3,4 or 5-phenylproline, cis 3,4 or 5 phenylproline
Proline	P	D-Pro, L-I-thioazolidine-4- carboxylic acid, D- or L-1- oxazolidine-4-carboxylic acid

Serine	s	D-Ser, Thr, D-Thr, allo-Thr, Met, D-Met, Met(O), D-Met(O), Val, D-Val
Threonine	T	D-Thr, Ser, D-Ser, allo-Thr, Met, D-Met, Met(O) D-Met(O), Val, D-Val
Tyrosine	Y	D-Tyr,Phe, D-Phe, L-Dopa, His,D-His
Valine	V	D-Val, Leu, D-Leu, Ile, D-Ile, Met, D-Met

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Other variants within the invention are those with modifications which increase peptide stability. Such variants may contain, for example, one or more nonpeptide bonds (which replace the peptide bonds) in the peptide sequence. Also included are: variants that include residues other than naturally occurring L-amino acids, such as D-amino acids or non-naturally occurring or synthetic amino acids such as beta or gamma amino acids and cyclic variants. Incorporation of D- instead of L-amino acids into the polypeptide may increase its resistance to proteases. See, e.g., U.S. 5,219,990.

The peptides of this invention may also be modified by various changes such as insertions, deletions and substitutions, either conservative or nonconservative where such changes might provide for certain advantages in their use.

In other embodiments, variants with amino acid substitutions which are less conservative may also result in desired derivatives, e.g., by causing changes in charge, conformation and other biological properties. Such substitutions would include for example, substitution of hydrophilic residue for a hydrophobic

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residue, substitution of a cysteine or proline for another residue, substitution of a residue having a small side chain for a residue having a bulky side chain or substitution of a residue having a net positive charge for a residue having a net negative charge. When the result of a given substitution cannot be predicted with certainty, the derivatives may be readily assayed according to the methods disclosed herein to determine the presence or absence of the desired characteristics.

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Variants within the scope of the invention include proteins and peptides with amino acid sequences having at least eighty percent homology with the extracellular region of CD40 or the extracellular region of CD40 ligand. More preferably the sequence homology is at least ninety percent, or at least ninety-five percent.

Just as it is possible to replace substituents of the scaffold, it is also possible to substitute functional groups which decorate the scaffold with characterized by similar features. These substitutions will initially be conservative, i.e., the replacement group will have approximately the same size, shape, hydrophobicity and charge as the original group. Nonsequence modifications may include, for example, in vivo or in vitro chemical derivatization of portions of naturally occurring CD40 or CD40 ligand, as well as changes in acetylation, methylation, phosphorylation, carboxylation or glycolsylation.

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In a further embodiment the protein, including the extracellular region of CD40 ligand and CD40, is modified by chemical modifications in which activity is preserved. For example, the proteins may be amidated, sulfated, singly or multiply halogenated, alkylated, carboxylated, or phosphorylated. The protein may also be singly or multiply acylated, such as with an acetyl group, with a

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farnesyl moiety, or with a fatty acid, which may be saturated, monounsaturat d or polyunsaturated. The fatty acid may also be singly or multiply fluorinated. invention also includes methionine analogs of the example the methionine and sulfone protein, for methionine sulfoxide analogs. The invention also includes salts of the proteins, such as ammonium salts, including alkyl or aryl ammonium salts, sulfate, hydrogen sulfate, phosphate, hydrogen phosphate, dihydrogen phosphate, thiosulfate, carbonate, bicarbonate, benzoate, sulfonate, thiosulfonate, mesylate, ethyl sulfonate and benzensulfonate salts.

The soluble, monomeric CD40-L protein can comprise all or part of the extracellular region of CD40-L. The extracellular region of CD40-L contains the domain that binds to CD40. Thus, soluble CD40-L can inhibit the interaction between CD40L and the CD40-bearing cell. This invention contemplates that sCD40-L may constitute the entire extracellular region of CD40-L, or a fragment or derivative containing the domain that binds to CD40.

Soluble CD40 protein (sCD40) comprises the extracellular region of CD40. sCD40 inhibits the interaction between CD40L and CD40-bearing cells. sCD40 may be in monomeric or oligomeric form.

In another embodiment of this invention the protein comprising soluble extracellular region of CD40 or portion thereof further comprises an Fc region fused to the extracellular region of CD40 or portion thereof. In a specific embodiment the Fc region is capable of binding to protein A or protein G. In another embodiment the Fc region comprises IgG, IgG₁, IgG₂, IgG₃, IgG₄, IgA, IgA₁, IgA₂, IgM, IgD, or IgE.

The soluble CD40/Fc fusion protein can be prepared using

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conventional techniques of enzymes cutting and ligation of fragments from desired sequences. Suitable Fc regions for the fusion protein are Fc regions that can bind to protein A or protein G, or that are capable of recognition by an antibody that can be used in purification or detection of a fusion protein comprising the Fc region. For example, the Fc region may include the Fc region of human IgG₁ or murine IgG₁. This invention also provides a nucleic acid molecule which encodes the CD40/Fc fusion protein.

The method of creating soluble forms of membrane molecules by recombinant means, in which sequences encoding the transmembrane and cytoplasmic domains are deleted, is well known. See generally Hammonds et al., U.S. Patent No. 5,057,417. In addition, methods of preparing sCD40 and CD40/Fc fusion protein are well-known. See, e.g., PCT International Publication No. WO 93/08207; Fanslow et al., "Soluble Forms of CD40 Inhibit Biologic Responses of Human B Cells, "J. Immunol., vol. 149, pp.655-60 (July 1992).

In an embodiment of this invention, the agent is selected by a screening method.

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In a specific embodiment the agent is selected by a screening method, which comprises isolating a sample of cells; culturing the sample under conditions permitting activation of CD40-bearing cells; contacting the sample with cells expressing a protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession no. HB 10916, or with a protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession no. HB 10916, effective to activate the CD40-bearing cells; contacting the sample with an amount of the agent effective to inhibit activation of the CD40-

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bearing cells if the agent is capable of inhibiting activation of the CD40-bearing cells; and determining whether the cells expressing the protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession no. HB 10916, or with the protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession no. HB 10916, activate the CD40-bearing cells in the presence of the agent. The cell sample may be isolated from diverse tissues, including cell lines in culture or cells isolated from an animal, such as dispersed cells from a solid tissue, cells derived from a bone marrow biopsy, or cells isolated from a body fluid such as blood or lymphatic fluid.

In another specific embodiment the agent (molecule) is selected based on a three-dimensional structure of soluble extracellular region of CD40 ligand or portion thereof capable of inhibiting interaction between CD40 ligand and CD40 on the cells. The agent may be selected from a library of known agents, modified from a known agent based on the three-dimensional structure, or designed and synthesized de novo based on the threedimensional structure. In specific embodiments the agent (molecule) is designed by structure optimization of a inhibitory agent based on a three-dimensional structure of a complex of the soluble extracellular region of CD40 ligand or portion thereof with the lead inhibitory agent. A lead inhibitory agent is a molecule which has been identified which, when it is contacted with CD40 ligand, binds to and complexes with the soluble extracellular region of CD40 ligand, CD40, or portion thereof, thereby decreasing the ability of the complexed or bound CD40 ligand or CD40 ligand portion to activate In another embodiment, a lead CD40-b aring cells. inhibitory agent may act by interacting with either the

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extracellular region of CD40 ligand, CD40, or in a tertiary complex with both a portion of CD40 ligand and CD40, decreasing the ability of the complexed CD40 ligand-CD40 to activate the CD40-bearing cells. In the methods of the invention, the CD40 ligand may be either soluble or bound to cells such as activated T cells, and may be either full length native CD40 ligand or portions Decreased ability to activate CD40-bearing cells may be measured in different ways. One way it may be measured is by showing that CD40 ligand, in the presence of inhibitor, causes a lesser degree activation of CD40-bearing cells, as compared treatment of the cells with a similar amount of CD40 ligand without inhibitor under similar conditions. Decreased ability to activate CD40-bearing cells may also be indicated by a higher concentration of inhibitor-CD40 ligand complex being required to produce a similar degree activation of CD40-bearing cells under similar conditions, as compared to unbound CD40 ligand. At the extreme, the inhibitor-contacted CD40 ligand may be unable to activate CD40-bearing cells at concentrations and under conditions which allow activation of these cells by unbound CD40 ligand or a given portion thereof.

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The agent (molecule) can be selected by a computational screening method using the crystal structure of a soluble fragment of the extracellular domain of human CD40L containing residues Gly116-Leu261 (sCD40L(116-261)).

The crystal structure to be used with the screening method has been determined at 2 Å resolution by the method of molecular replacement. In brief, a soluble fragment of the extracellular domain of human CD40 ligand containing amino acid residues Gly 116 to the c-terminal residue Leu 261 was first produced in soluble form, then purified and crystalliz d. The crystals were used to collect diffraction data. Molecular replacement and

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refinement were done with the XPLOR program package and QUANTA (Molecular Simulations, Inc.) Software. particular, a 3-dimensional model of human sCD40L was constructed using the murine CD40L model using QUANTA protein homology modeling software. This model was used as a probe for crystallographic analysis calculations and This method of determining the refined using XPLOR. crystal structure of sCD40L is described in more detail in Karpusas et al., "2 Å crystal structure of an extracellular fragment of human CD40 ligand," Structure The atomic coordinates (October 1995) 3(10):1031-1039. of sCD40L(116-261) are provided in Figures 1A-Y. The for selecting an agent screening method iterative structure drug design and computational optimization, as described below.

selected using inhibitor be an The agent may computational drug design. Using this method, the sCD40L crystal structure coordinates are used as an input for a computer program, such as DOCK, which outputs a list of 20 molecular structures that are expected to bind to CD40L. Use of such computer programs is well-known. Kuntz, "Structure-Based Strategies for drug design and discovery," Science, vol. 257, p. 1078 (1992). The list structures can then screened by be molecular 25 biochemical assays for CD40L binding. Competition-type biochemical assays, which are well known, can be used. See, e.g., Bajorath et al., "Identification of residues of CD40 and its ligand which are critical for the receptorligand interaction," Biochemistry, 34, p. 1833 (1995). 30 The structures that are found to bind to CD40L can thus be used as agents for the present invention. The agent may also be a modified or designed molecule, determined by interactive cycles of structure optimization. this approach, a small molecule inhibitor of CD40L found 35 using the ab v computational approach or other approach can be co-crystallized with sCD40L and the crystal

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structure of the complex solved by molecular replacement. The information revealed through molecular replacement can b used to optimize the structure of the inhibitors by clarifying how the molecules interact with CD40L. The molecule may be modified to improve its physiochemical properties, including specificity and affinity for CD40L.

In an embodiment of this invention the agent is a small molecule. As used herein a small molecule is a compound having a molecular weight between 20 Da and 1x10⁶ Da, preferably from 50 Da to 2 kDa.

This invention also provides a method of inhibiting activation by CD40 ligand of renal cells bearing CD40 on the surface of the cells, in a subject, comprising administering to the subject an agent capable of inhibiting interaction between CD40 ligand and CD40 on the cells, the agent being present in an amount effective to inhibit activation of the cells in the subject.

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In specific embodiments the CD40-bearing renal cells are selected from the group consisting of glomerular endothelial cells, mesangial cells, distal tubules, proximal tubules, parietal epithelial cells, visceral epithelial cells, cells of a Henle loop or limb thereof, and interstitial inflammatory cells. In a more specific embodiment the parietal epithelial cells are crescent parietal epithelial cells.

In an embodiment of this invention the agent inhibits binding of CD40 ligand to CD40 on the cells.

In an embodiment of this invention the agent is a protein. In another embodiment of this invention the agent is a nonprotein.

In a specific embodiment the protein comprises an

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antibody or portion thereof capable of inhibiting any interaction between CD40 ligand and CD40 on the cells. The antibody is a monoclonal or polyclonal antibody. In a more specific embodiment the monoclonal antibody specifically binds to the epitope to which monoclonal antibody 5c8 (ATCC Accession No. HB 10916) specifically binds. An example of such a monoclonal antibody is monoclonal antibody 5c8 (ATCC Accession No. HB 10916). In other embodiments the monoclonal antibody is a chimeric antibody or a humanized antibody.

In a specific embodiment the portion of the antibody comprises a complementarity determining region or variable region of a light or heavy chain. In another specific embodiment the portion of the antibody comprises a complementarity determining region or a variable region. In another specific embodiment the portion of the antibody comprises a Fab or a single chain antibody.

In another embodiment the protein comprises soluble extracellular region of CD40 ligand or portion thereof capable of inhibiting any interaction between CD40 ligand and CD40 on the cells; or soluble extracellular region of CD40 or portion thereof capable of inhibiting any interaction between CD40 ligand and CD40 on the cells. In a specific embodiment the soluble extracellular region of CD40 ligand or CD40 is a monomer. In another embodiment the soluble extracellular region of CD40 is an oligomer.

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In another embodiment of this invention the protein comprising soluble extracellular region of CD40 or portion thereof further comprises an Fc region fused to the extracellular region of CD40 or portion thereof. In a specific embodiment the Fc region is capable of binding to protein A or protein G. In another specific embodiment the Fc region comprises IgG, IgG₁, IgG₂, IgG₃,

IgG,, IgA, IgA,, IgA,, IgM, IgD, or IgE.

The subject which can be treated by the above-described methods is an animal. Preferably the animal is a mammal. Examples of mammals which may be treated include, but are not limited to, humans, non-human primates, rodents (including rats, mice, hamsters and guinea pigs) cow, horse, sheep, goat, pig, dog and cat.

10 In an embodiment of this invention, the agent is selected by a screening method.

In a specific embodiment the agent is selected by a screening method, which comprises isolating a sample of 15 cells; culturing the sample under conditions permitting activation of CD40-bearing cells; contacting the sample with cells expressing a protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession no. HB 10916, or with a 20 protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession no. HB 10916, effective to activate the CD40bearing cells; contacting the sample with an amount of the agent effective to inhibit activation of the CD40-25 bearing cells if the agent is capable of inhibiting activation of the CD40-bearing cells; and determining whether the cells expressing the protein which specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession no. HB 30 or with the protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession no. HB 10916, activate the CD40-bearing cells in the presence of the agent. cell sample may be isolated from diverse tissues, 35 including cell lines in culture or cells isolated from an animal, such as dispersed cells from a solid tissue, cells derived from a bone marrow biopsy, or cells

isolated from a body fluid such as blood or lymphatic fluid.

In another specific embodiment the molecule (agent) is selected based on a three-dimensional structure of soluble extracellular region of CD40 ligand or portion thereof capable of inhibiting any interaction between CD40 ligand and CD40 on the cells. The molecule may be selected from a library of known molecules, modified from the three-dimensional known molecule based on 10 structure, or designed and synthesized de novo based on the three-dimensional structure. In specific embodiments agent or molecule is designed by optimization of a lead inhibitory agent based on a threedimensional structure of a complex of the soluble 15 extracellular region of CD40 ligand or portion thereof with the lead inhibitory agent.

Method of Treatment

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This invention provides a method of treating, in a subject, an inflammatory kidney disease, comprising the above-described method of inhibiting activation by CD40 ligand of renal cells bearing CD40 on the surface of the cells, which comprises administering to the subject an agent capable of inhibiting interaction between CD40 ligand and CD40 on the cells, the agent being present in an amount effective to inhibit activation of the cells in the subject, thereby treating the inflammatory kidney disease.

The inflammatory kidney disease may be one which is initiated by autoantibody deposition in kidney, or one which is not initiated by autoantibody deposition in kidney. Many kidney diseases for which the methods of the invention are useful include ones which have multifactorial etiology.

In an embodiment of this invention the kidney disease is sel cted from the group consisting of: membranous glomerulonephritis, minimal change disease/acute tubular pauci-immune glomerulonephritis; necrosis; segmental glomerulosclerosis; interstitial nephritis; 5 antitissue antibody-induced glomerular injury, such as anti-basement membrane antibody disease; circulating immune-complex disease; glomerulopathies associated with multisystem diseases; drug-induced glomerular disease; 10 renal transplant rejection; rapidly progressive glomerulonephritis; and post-streptococcal glomerulonephritis. Circulating immune-complex diseases infective endocarditis, leprosy, syphilis, hepatitis B, malaria, and diseases of endogenous antigens 15 such as DNA, thyroglobulin, autologous immunoglobulins, erythrocyte stroma, renal tubule antigens, and tumorspecific or tumor-associated antigens. Glomerulopathies associated with multisystem diseases include diabetic nephropathy, systemic lupus erythematosus, Goodpasture's disease, vasculitis, multiple myeloma, 20 Waldenström's macroglobulinemia, and amyloidosis. In specific embodiments the vasculitis is Henoch-Schönlein purpura, polyarteritis nodosa (sometimes called polyarteritis), Wegener's granulomatosis, cryoglobulinemia (sometimes 25 called cryoimmunoglobulinemia). The kidney disease may also be one which affects the renal tubules, such as toxins, neoplasias, hypersensitivity nephropathy, Sjögren's syndrome, and AIDS. In a specific embodiment the pauci-immune glomerulonephritis is ANCA+ pauci-immune 30 glomerulonephritis, or Wegener's granulomatosis. another specific embodiment the interstitial nephritis is drug-induced interstitial nephritis.

The compounds of this invention may be administered in any manner which is medically acceptable. This may include injections, by parenteral routes such as intravenous, intravascular, intraarterial, subcutaneous,

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intramuscular, intratumor, intraperitoneal, intraventricular, intraepidural, or others as well as oral, nasal, ophthalmic, rectal, topical, or inhaled. Sustained release administration is also specifically included in the invention, by such means as depot injections of erodible implants directly applied during surgery.

The compounds are administered at any dose per body weight and any dosage frequency which is medically 10 Acceptable dosage includes a range of acceptable. between about 0.01 and 200 mg/kg subject body weight. A preferred dosage range is between about 0.1 and 50 mg/kg. Particularly preferred is a dose of between about 1 and The dosage is repeated at intervals ranging 15 30 mg/kg. from each day to every other month. One preferred dosing regimen is to administer a compound of the invention daily for the first three days of treatment, after which the compound is administered every 3 weeks, with each administration being intravenously at 5 or 10 mg/kg body 20 Another preferred regime is to administer a compound of the invention daily intravenously at 5 mg/kg body weight for the first three days of treatment, after which the compound is administered subcutaneously or intramuscularly every week at 10 mg per subject. Another . 25 preferred regime is to administer a single dose of the compound of the invention parenterally at 20 mg/kg body followed by administration of the compound subcutaneously or intramuscularly every week at 10 mg per subject. 30

The compounds of the invention may be administered as a single dosage for certain indications such as preventing immune response to an antigen to which a subject is exposed for a brief time, such as an exogenous antigen administered on a single day of treatment. Examples of such an antigen would include coadministration of a

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compound of the invention along with a gene therapy vector, or a therapeutic agent such as an antigenic pharmaceutical or a blood product. In indications where antigen is chronically present, such as in controlling immune reaction to transplanted tissue or to chronically administered antigenic pharmaceuticals, the compounds of the invention are administered at intervals for as long a time as medically indicated, ranging from days or weeks to the life of the subject.

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Inflammatory responses are characterized by redness, swelling, heat and pain, as consequences of capillary dilation with edema and migration of phagocytic leukocytes. Inflammation is further defined by Gallin (Chapter 26, Fundamental Immunology, 2d Ed., Raven Press, York, 1989, pp. 721-733), which is herein incorporated by reference.

This invention will be better understood from the Experimental Details which follow. However, one skilled in the art will readily appreciate that the specific methods and results discussed are merely illustrative of the invention as described more fully in the claims which follow thereafter.

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Experimental Details

CD40 expression in normal kidney and in renal biopsy specimens obtained from patients with systemic lupus erythematosus and other kidney diseases was examined.

Patients and Methods

Immunohistochemistry

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Immunohistochemical analyses of frozen sections were performed with a V ctastain Elite Kit (Vector,

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Burlingame, CA) as previously described. Briefly, the tissue was first blocked with PBS containing horse serum and 1% BSA and additional blocking was obtained utilizing an Avidin/Biotin Blocking Kit also purchased from Vector. The sections were then stained with 1:1000 dilutions of anti-CD40 mAb G28.5 or an isotype control mAb in PBS biotinylated horse anti-mouse followed by Endogenous peroxidase activity was blocked with 1:400 dilution of H_2O_2 . Bound antibody was visualized with the Vectastain ABC reagent followed by the chromogen 3-amino-9-ethylcarbazole (Vector Laboratories). The tissue was counterstained with Mayer's hematoxylin (Sigma).

Staining was evaluated visually. In the following tables "O" indicates no staining; 1+ indicates minimal staining; 2+ indicates moderate staining; and 3+ indicates intense staining.

Results

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Analysis of CD40 expression in normal kidney

Initial studies of renal CD40 expression were prompted by the observation that CD40 is normally expressed on endothelial cells in a variety of tissues. Consistent with this finding, it was found that renal interstitial capillaries and larger vessels express CD40. also found to be expressed on other renal parenchymal cells, such as glomerular endothelial cells, glomerular mesangial cells and parietal epithelial cells of Bowman's Glomerular visceral epithelial cells do not Distal tubules are strongly immunoreactive express CD40. for CD40 and staining was most intense along the basolateral membrane. In contrast, proximal tubules are not immunoreactive with anti-CD40 mAb. An isotype c ntrol mAb did not stain renal specimens. immunoreactivity noted with anti-CD40 mAb G28.5 is most

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likely specific and does not represent cross-reactivity becaus similar staining was noted with an additional anti-CD40 mAb. Thus it is concluded that renal parenchymal cells differentially express CD40.

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Analysis of renal CD40 expression in systemic lupus erythematosus

- Whether renal CD40 expression is upregulated in lupus glomerulonephritis was analyzed. Frozen sections obtained from patient biopsy specimens were stained with anti-CD40 mAb G28.5 or an isotype control mAb.
- 15 Renal CD40 expression in systemic lupus erythematosus was analyzed. Patients with Class III and IV lupus nephritis tended to have increased CD40 expression on glomerular endothelial cells, mesangial cells and distal tubules. In addition, proximal tubules are CD40+ in patients with 20 Class III and IV lupus nephritis. Also, there is striking CD40 expression on parietal epithelial cells in patients with crescent formation. CD40 is also present on interstitial inflammatory cells. The distribution and intensity of renal CD40 expression in patients with pure 25 Class V disease was similar to that seen in normal kidney.

Whether renal CD40 upregulation was unique to systemic lupus erythematosus was investigated. To do so, CD40 expression was investigated in patients with the following renal diseases: membranous glomerulonephritis, minimal change disease/acute tubular necrosis, ANCA+ pauci-immune glomerulonephritis, focal glomerulosclerosis and IqA nephropathy. Proximal tubule CD40 expression was upregulated in ANCA+ pauci-immune glomerulonephritis, focal segmental glomerulosclerosis In contrast, there was little and IgA nephropathy.

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proximal tubule CD40 immunoreactivity in membranous glomerulonephritis r minimal change disease/acute tubular necrosis. Crescent parietal epithelial cells in IgA nephropathy are also striking CD40+. Interstitial inflammatory cells, when present, also express CD40. These findings demonstrate that CD40 expression is upregulated in a variety of inflammatory renal diseases. Moreover, these studies indicate that CD40L mediated interactions with renal parenchymal cells play roles in normal renal physiology and augment inflammatory responses in renal diseases.

Table 2: CD40 Expression In Normal Kidney

		Glomerular	lar		Inters	Interstitium		Tubules	
Sp cimen	EC	Mesangial	VEC	PEC	Cap EC	Leukocytes Proximal	Proximal	Distal	Collecting
-	1+diffuse	1+	,	-,;	2+diffuse	,		- 1	
								STUTTORS	+7
2	1+	2+diffuse	1	1+focal	1+focal 2+diffuse	1+(rare)	1	3+diffuse	3+
3									

Table 3: CD40 Expression In SLE Glomerulonephritis Interstitium

les	Distal	2+	3+	2+	3+	.2+	3+	3+	3+	3+	3+	3+	3+	3+	3+	2+	2	2+	1+	
Tubules	Proximal	-	2+	1+	2-3+	1+	2+	2-3+	3+	2+	3+	1+	1-2+	2+	3+	0	+1	t	1+	
Interstitium	Leukocytes	0	3+	1+	3+	1+	2+	2+	3+	3+	2+	3+	3+	1+	3+	1+	1	No leuk	1+	
Inters	Cap EC	2+	+1	1+	1+	1+	1+	1+	. 1+	1+	2+	3+	2+	2+	2+	2+	+1	1+	1+	
Table 3: Cra	WHO Class	lib	III	V/III	V/VI-III	ΛI	IV	ΙΛ	IV	IV	ΛI	VI	IV/VI	IV/V	IV/V	Λ	Λ	۸	Λ	,
	Patient	KC95-94	KC95-277	KC95-286	KC94-78	KC95-308	KC94-269	K94-165	K94-59	K95-089	K94-6	K94-12	K95-090	K95-003	KC95-264	K95-7	KC95-195	K94-142	K95-12	N2J 14

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Table 4: CD40 Expression In SLE Glomerulonephritis

	PEC	+1	1+	1+	1+	1+	3+	3+	+1	sclero	1+	3+ (cresc)	0	+1	-(par 1+)	0(par 1+)
	VEC	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Expression	Mesangial	1+	3+	2+	3+	2-3+	3+	2+	3+	0	1+	3+	1+	+1	1	1
Glomerular	EC	0	3+	2+	3+	1+	.3+	3+	3+	1+	2+	3+	1+	. 1	0	0
	WHO Class	III	V/VI-III	IV	IV	IV	ΛI	ΛI	IV	1//1	IV/V	V/VI	Λ	Λ	Λ	Λ
	Patient	KC95-277	KC94-78	KC95-308	KC94-269	K94-165	K94-59	K95-089	K94-12	K95-090	K95-003	KC95-264	K95-7	KC95-195	K94-142	K95-12

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Table 5: CD40 Expression In Non-SLE Glomerulonephritis

Glomerular Expression

			744	730
Renal Disease LC	ည ချ	Mesangiai	VEC	797
		-		
Membranous 0	0	1+	0	1+
MC/ATN 0	0	+1	0	1+
Pauci-immune 1	1	.1	0	
FSGS 0	0	+	0	
IgA 1+	1+	2+	0	2+
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Table 6: CD40 Expression In Non-SLE Glomerulonephritis

				,	,	,
Distal		2+	1+	3+	3+	3+
Proximal		1+	0	2+	2+	2+
Leukocytes		+1	2+	3+	3+	3+
Cap EC		1+	+1	1+	1+	1+
Renal Disease		Membranous	MC/AIN	Pauci-immune	FSGS	IgA
Patient		KC95-310	KC95-299	KC95-312	KC95-280	KC94-282
	Renal Disease Cap EC Leukocytes Proximal	Renal Disease Cap EC Leukocytes Proximal	Renal DiseaseCap ECLeukocytesProximalMembranous1+±1+	Renal Disease Cap EC Leukocytes Proximal Membranous 1+ ± 1+ MC/ATN ± 0	Renal Disease Cap EC Leukocytes Proximal Membranous 1+ ± 1+ MC/ATN ± 0 0 Pauci-immune 1+ 3+ 2+	Renal Disease Cap EC Leukocytes Proximal Membranous 1+ ± 1+ MC/ATN ± 2+ 0 Pauci-immune 1+ 3+ 2+ FSGS 1+ 3+ 2+

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Analysis of renal CD40-ligand expressi n in inflammatory renal dis as s

In situ CD40L expression was studied in renal biopsy specimens from patients with SLE GN (n=18), as well as in normal kidney and biopsy specimens from patients with IgA nephropathy, focal segmental glomerulosclerosis, minimal change disease, idiopathic membranous GN and ANCA pauciimmune GN. Immunohistochemical studies were performed on frozen sections utilizing anti-CD40L mAb 5C8 or controls mAbs. Upregulation of CD40L expression is observed in class IV lupus glomerulonephritis (Figures 4A, 4B and 5), focal segmental glomerulosclerosis (Figure 7) and Iga nephropathy (Figure 9). CD40L expression is noted as dim, discrete staining of some infiltrating mononuclear cells. These results provide further evidence that CD40L mediated signals play a role in the immunopathogenesis of inflammatory glomerular or tubulointerstitial diseases by interacting with CD40 target cells in the kidney.

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANTS: Yellin, Michael J.

Lederman, Seth Chess, Leonard Karpusas, Mihail N. Thomas, David W.

(ii) TITLE OF INVENTION: THERAPEUTIC APPLICATIONS OF T-BAM

(CD40-L) TECHNOLOGY TO TREAT

INFLAMMATORY KIDNEY DISEASES

- (iii) NUMBER OF SEQUENCES: 1
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Cooper & Dunham LLP
 - (B) STREET: 1185 Avenue of the Americas
 - (C) CITY: New York
 - (D) STATE: New York
 - (E) COUNTRY: USA
 - (F) ZIP: 10036
 - (V) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: Not Yet Known
 - (B) FILING DATE: Herewith
 - (C) CLASSIFICATION:
- (vii) PREVIOUS APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/587,334
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 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: White Esq., John P.
 - (B) REGISTRATION NUMBER: 28,678
 - (C) REFERENCE/DOCKET NUMBER: 48558-B
 - (ix) TELECOMMUNICATION INFORMATION:
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 - (B) TELEFAX: (212)391 0525
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 146 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein

(iii) HYPOTHETICAL: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

Gly Asp Gln Asn Pro Gln Ile Ala Ala His Val Ile Ser Glu Ala Ser 1 5 10 15

Ser Lys Thr Thr Ser Val Leu Gln Trp Ala Glu Lys Gly Tyr Tyr Thr 20 25 30

Met Ser Asn Asn Leu Val Thr Leu Glu Asn Gly Lys Gln Leu Thr Val 35 40 45

Lys Arg Gln Gly Leu Tyr Tyr Ile Tyr Ala Gln Val Thr Phe Cys Ser 50 55 60

Asn Arg Glu Ala Ser Ser Gln Ala Pro Phe Ile Ala Ser Leu Cys Leu 65 70 75 80

Lys Ser Pro Gly Arg Phe Glu Arg Ile Leu Leu Arg Ala Ala Asn Thr 85 90 95

His Ser Ser Ala Lys Pro Cys Gly Gln Gln Ser Ile His Leu Gly Gly 100 105 110

Val Phe Glu Leu Gln Pro Gly Ala Ser Val Phe Val Asn Val Thr Asp 115 120 125

Pro Ser Gln Val Ser His Gly Thr Gly Phe Thr Ser Phe Gly Leu Leu 130 135 140

Lys Leu 145

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What is claim d is:

 A method of inhibiting activation by CD40 ligand of renal cells bearing CD40 on the surface of the cells, comprising contacting the cells with an agent capable of inhibiting interaction between CD40 ligand and CD40 on the cells, the agent being present in an amount effective to inhibit activation of the cells.

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- 2. The method of claim 1, wherein the CD40-bearing renal cells are selected from the group consisting of glomerular endothelial cells, mesangial cells, distal tubule cells, proximal tubule cells, parietal epithelial cells, visceral epithelial cells, cells of a Henle limb, and interstitial inflammatory cells.
- 3. The method of claim 2, wherein the parietal epithelial cells are crescent parietal epithelial cells.
 - 4. The method of claim 1, wherein the agent inhibits binding of CD40 ligand to CD40 on the cells.

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- 5. The method of claim 1, wherein the agent is a protein.
- 6. The method of claim 5, wherein the protein comprises an antibody or portion thereof.
 - 7. The method of claim 6, wherein the antibody is a monoclonal antibody.
- 35 8. The method of claim 7, wherein the monoclonal antibody specifically binds to the epitope to which monoclonal antibody 5c8 (ATCC Accession No. HB

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10916) specifically binds.

- 9. The method of claim 8, wherein the monoclonal antibody is monoclonal antibody 5c8 (ATCC Accession No. HB 10916).
 - 10. The method of claim 7, wherein the monoclonal antibody specifically binds to CD40.
- 10 11. The method of claim 10, wherein the antibody is humanized, chimeric, or primatized.
 - 12. The method of claim 7, wherein the monoclonal antibody is a chimeric antibody.
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 13. The method of claim 7, wherein the monoclonal antibody is a humanized antibody.
- 14. The method of claim 6, wherein the portion of the 20 antibody comprises a complementarity determining region or variable region of a light or heavy chain.
- 15. The method of claim 6, wherein the portion of the antibody comprises a complementarity determining25 region or a variable region.
 - 16. The method of claim 15, wherein the portion of the antibody comprises a Fab or a single chain antibody.
- 30 17. The method of claim 5, wherein the protein comprises soluble extracellular region of CD40 ligand, or variant thereof including conservative substituents, or portion thereof; or soluble extracellular region of CD40, or variant thereof including conservative substituents, or portion thereof.
 - 18. The method of claim 17, wherein the soluble

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extracellular region of CD40 ligand or CD40 is a monomer.

- 19. The method of claim 17, wherein the soluble extracellular region of CD40 is an oligomer.
- 20. The method of claim 17, wherein the protein comprising soluble extracellular region of CD40 or portion thereof or CD40 ligand or portion thereof further comprises an Fc region fused to the extracellular region of CD40 or portion thereof or CD40 ligand or portion thereof.
- 21. The method of claim 20, wherein the Fc region is capable of binding to protein A or protein G.
 - 22. The method of claim 21, wherein the Fc region comprises IgG, IgA, IgM, IgD, or IgE, or subclasses thereof.

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- 23. The method of claim 22, wherein: the IgG is IgG₁, IgG₂, IgG₃, or IgG₄; or the IgA is IgA₁ or IgA₂.
- 25 24. The method of claim 1, wherein the agent is nonprotein.
 - 25. The method of claim 1, wherein the agent is selected from a library of known agents.

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- 26. The method of claim 1, wherein the agent is modified from a known agent.
- 27. The method of claim 26, wherein the modified agent is designed by structure optimization of a lead inhibitory agent based on a three-dimensional structure of a complex of soluble extracellular

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region of CD40 ligand or portion thereof with the lead inhibitory agent.

5 28. The method of claim 1, wherein the agent is selected by a screening method, which comprises:

isolating a sample of cells;

culturing the sample under conditions permitting activation of CD40-bearing cells;

contacting the sample with cells expressing a protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession No. HB 10916, or with a protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession No. HB 10916, effective to activate the CD40-bearing cells;

contacting the sample with an amount of the agent effective to inhibit activation of the CD40-bearing cells if the agent is capable of inhibiting activation of the CD40-bearing cells; and

determining whether the cells expressing the protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession No. HB 10916, or with the protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession No. HB 10916, activate the CD40-bearing cells in the presence of the agent.

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29. The method of claim 28, wherein the agent is

selected from a library of known agents.

30. The method of claim 29, wherein the known agents are nonprotein agents.

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- 31. A method of inhibiting activation by CD40 ligand of renal cells bearing CD40 on the surface of the cells, in a subject, comprising administering to the subject an agent capable of inhibiting interaction between CD40 ligand and CD40 on the cells, the agent being present in an amount effective to inhibit activation of the cells in the subject.
- 32. The method of claim 31, wherein the CD40-bearing renal cells are selected from the group consisting of glomerular endothelial cells, mesangial cells, distal tubule cells, proximal tubule cells, parietal epithelial cells, visceral epithelial cells, cells of a Henle limb, and interstitial inflammatory cells.
 - 33. The method of claim 32, wherein the parietal epithelial cells are crescent parietal epithelial cells.

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- 34. The method of claim 31, wherein the agent inhibits binding of CD40 ligand to CD40 on the cells.
- 35. The method of claim 31, wherein the agent is a protein.
 - 36. The method of claim 35, wherein the protein comprises an antibody or portion thereof.
- 35 37. The method of claim 36, wherein the antibody is a monoclonal antibody.

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38. The method of claim 37, wherein the monoclonal antibody specifically binds to the epitope to which monoclonal antibody 5c8 (ATCC Accession No. HB 10916) specifically binds.

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- 39. The method of claim 38, wherein the agent is monoclonal antibody 5c8 (ATCC Accession No. HB 10916).
- 10 40. The method of claim 37, wherein the monoclonal antibody specifically binds to CD40.
 - 41. The method of claim 40, wherein the antibody is humanized, chimeric, or primatized.

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- 42. The method of claim 37, wherein the monoclonal antibody is a chimeric antibody.
- 43. The method of claim 37, wherein the monoclonal antibody is a humanized antibody.
 - 44. The method of claim 36, wherein the portion of the antibody comprises a complementarity determining region or variable region of a light or heavy chain.

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- 45. The method of claim 36, wherein the portion of the antibody comprises a complementarity determining region or a variable region.
- 30 46. The method of claim 45, wherein the portion of the antibody comprises a Fab or a single chain antibody.
 - 47. The method of claim 31, wherein the subject is a mammal.

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48. The method of claim 47, wherein the mammal is a rodent.

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49. The method of claim 47, wherein the mammal is a human.

- 50. The method of claim 31, wherein the protein comprises soluble extracellular region of CD40 ligand, or variant thereof including conservative substituents, or portion thereof; or soluble extracellular region of CD40, or variant thereof including conservative substituents, or portion thereof.
 - 51. The method of claim 50, wherein the soluble extracellular region of CD40 ligand or CD40 is a monomer.

52. The method of claim 50, wherein the soluble extracellular region of CD40 is an oligomer.

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- 53. The method of claim 50, wherein the protein comprising soluble extracellular region of CD40 or portion thereof or CD40 ligand or portion thereof further comprises an Fc region fused to the extracellular region of CD40 or portion thereof or CD40 ligand or portion thereof.
 - 54. The method of claim 53, wherein the Fc region is capable of binding to protein A or protein G.
- 55. The method of claim 53, wherein the Fc region comprises IgG, IgA, IgM, IgD, or IgE, or subclasses thereof.
 - 56. The method of claim 55, wherein:
 the IgG is IgG₁, IgG₂, IgG₃, or IgG₄; or
 the IgA is IgA₁ or IgA₂.
 - 57. The method of claim 31, wherein the agent is

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nonprotein.

- 58. The method of claim 57, wherein the agent is a small molecule.
- 59. The method of claim 31, wherein the agent is selected from a library of known agents.
- 60. The method of claim 31, wherein the agent is modified from a known agent.
 - 61. The method of claim 60, wherein the modified agent is designed by structure optimization of a lead inhibitor based on a three-dimensional structure of a complex of soluble extracellular region of CD40 ligand or portion thereof with the lead inhibitor.
 - 62. The method of claim 31, wherein the agent is selected by a screening method, which comprises:
 - isolating a sample of cells;

culturing the sample under conditions permitting activation of CD40-bearing cells;

contacting the sample with cells expressing a protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession No. HB 10916, or with a protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession No. HB 10916, effective to

contacting the sample with an amount of the agent effective to inhibit activation of the CD40-bearing cells if the agent is capable of inhibiting

activate the CD40-bearing cells;

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activation of the CD40-bearing cells; and

determining wh ther the cells expressing the protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession No. HB 10916, or with the protein which is specifically recognized by monoclonal antibody 5c8 produced by the hybridoma having ATCC Accession No. HB 10916, activate the CD40-bearing cells in the presence of the agent.

- 63. The method of claim 62, wherein the agent is selected from a library of known agents.
- 15 64. The method of claim 63, wherein the known agents are nonprotein agents.
- 65. A method of treating, in a subject, an inflammatory kidney disease, comprising inhibiting activation by CD40 ligand of renal cells bearing CD40 on the surface of the cells, according to the method of claim 31.
- 66. The method of claim 65, wherein the inflammatory kidney disease is not initiated by autoantibody deposition in kidney.
 - 67. The method of claim 65, wherein the kidney disease is selected from the group consisting of:
- membranous glomerulonephritis;
 minimal change disease/acute tubular necrosis;
 pauci-immune glomerulonephritis;
 focal segmental glomerulosclerosis;
 interstitial nephritis;
 antitissue antibody-induced glomerular injury;
- antitissue antibody-induced glomerular injury; circulating immune-c mplex disease; a glomerulopathy associated with a multisystem

disease:

drug-induc d glomerular disease;
renal transplant rejection;
rapidly progressive glomerulonephritis; and
post-streptococcal glomerulonephritis.

68. The method of claim 67, wherein the antitissue antibody-induced glomerular injury is anti-basement membrane antibody disease.

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69. The method of claim 67, wherein the circulating immune-complex disease is selected from the group consisting of:

infective endocarditis;

15 leprosy;

syphilis;

hepatitis B;

malaria; and

a disease associated with an endogenous antigen.

- 70. The method of claim 69, wherein the endogenous antigen is DNA, thyroglobulin, an autologous immunoglobulin, erythrocyte stroma, a renal tubule antigen, a tumor-specific antigen, or a tumor-associated antigen.
- 71. The method of claim 67 wherein the glomerulopathy associated with a multisystem disease is selected from the group consisting of:

diabetic nephropathy; systemic lupus erythematosus;

Goodpasture's disease;

vasculitis;

35 multiple myeloma;

Waldenström's macroglobulinemia; and amyloidosis.

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72. The method of claim 71, wherein the vasculitis is selected from the group consisting of:

Henoch-Schönlein purpura; polyarteritis nodosa; Wegener's granulomatosis; and cryoglobulinemia.

- 73. The method of claim 67, wherein the pauci-immune glomerulonephritis is ANCA+ pauci-immune glomerulonephritis, or Wegener's granulomatosis.
 - 74. The method of claim 67, wherein the interstitial nephritis is drug-induced interstitial nephritis.
- 15 75. The method of claim 65 wherein the kidney disease affects renal tubules.
- 76. The method of claim 75, wherein the kidney disease which affects renal tubules is selected from the group consisting of:

a kidney disease associated with a toxin; a neoplasia; hypersensitivity nephropathy; Sjögren's syndrome; and

25 AIDS.

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FIGURE 1A

				NATES	OF CD401 CRYSTAL STRUCTURE IN PDB FORMAT 90.460 90.00 90.00 120.00 R3
CHYST		.170	GLY	116	B AR. 16 A BR
ATOM	i		GLY	116	n 44n 45 45n na 44n
ATOM	2				
ATOM	3		GLY	116	
ATOM	4		GLY	116	
ATOM	5	CA	GLY	116	
ATOM	6	С	GLY	116	-6.990 -16.621 24.780 1.0C 64.34 A
ATOM	7	0	GLY	116	-6.968 -17.814 24.563 1.00 64.44 A
ATOM	8	N	ASP	117	
ATOM	9	H	ASP	117	-5.617 -16.709 26.170 1.00 15.00 A
ATOM	10	CA	ASP	117	
MOTA	11	CB	ASP	117	-5.711 -14.402 27.539 1.00 63.36 A
ATOM	12	CG	ASP	117	-6.518 -15.163 28.574 1.00 63.71 A
ATOM	13	OD1	ASP	117	-6.090 -16.247 28.96 5 1.00 63.24 A
ATOM	14	QD2	ASP	117	-7.566 -14.66B 28.987 1.00 63.29 A
ATOM	15	C	ASP	117	-5.651 -13.585 25.184 1.00 63.31 A
ATOM	16	0	ASP	117	-6.039 -12.427 25.145 1.00 63.35 A
ATOM	17	N	GLN	118	-4.713 -14.090 24.379 1.00 62.72 A
ATOM	18	Н	GLN	118	-4.450 -15.040 24.541 1.00 15.00 A
ATOM	19	CA	GLN	118	-4.097 -13.313 23.281 1.00 61.79 A
ATOM	20	CB	GLN	118	-2.918 -14.117 22.687 1.00 62.46 A
ATOM	21	CG	GLN	118	-3.047 -15.659 22.562 1.00 62.95 A
ATOM	22	CD	GLN	118	-4.277 -16.118 21.790 1.00 63.26 A
ATOM	23		GLN	118	-5.396 -16.000 22.277 1.00 63.43 A
ATOM	24		GLN	118	-4.044 -16.665 20.601 1.00 63.42 A
ATOM		HE21		118	-4.836 -16.715 19.975 1.00 15.00 A
ATOM	26	HE22		118	-3.151 -16.995 20.298 1.00 15.00 A
	27	C	GLN	118	
ATOM		Ö	GLN	118	
ATOM	28				_ iii iii iii iii iii iii iii iii iii i
ATOM	29	N	asn Asn	119 119	
ATOM	30	H			
ATOM	31	CA	ASN	119	
ATOM	32	CB	asn Asn	119	
ATOM	33	CG		119	-7.652 -13.352 20.375 1.00 57.45 A
ATOM	34	OD1		119	-7.941 -14.303 21.084 1.00 58.50 A
ATOM	35	ND2		119	-7.005 -13.431 19.241 1.00 58.58 A
ATOM		HD21		119	-6.843 -12.617 18.646 1.00 15.00 A
ATOM	37	HD22		119	-6.740 -14.221 18.684 1.00 15.00 A
ATOM	38	C	ASN	119	-7.053 -9.724 21.571 1.00 53.62 A
ATCM	39	0	ASN	119	-6.746 -8.933 20.694 1.00 56.55 A
ATOM	40	N	PRO	120	-7.737 -9.288 22.698 1.00 50.17 A
ATOM	41	CD	PRO	120	-8.151 -10.129 23.810 1.00 51.90 A
ATOM	42	CA	PRO	120	-8.402 -7.945 22.818 1.00 48.19 A
ATOM	43	CB	PRO	120	-9.191 -8.008 24.117 1.00 47.42 A
ATOM	44	CG	PRO	120	-9.444 -9.493 24.321 1.00 51.93 A
ATOM	45	С	PRJ	125	-7.750 -6.524 22.657 1.00 45.59 A
ATOM	46	C	PRO	120	-8.187 -5.516 23.225 1.00 45.37 A
ATOM	47	N	GLN	121	-6.789 -6.458 21.721 1.00 38.52 A
ATOM	48	H	GLN	121	-6.287 -7.304 21.505 1.00 15.00 A
ATOM	49	CA	GLN	121	-6.733 -5.359 20.753 1.00 29.14 A
ATOM	50	CB	GLN	121	-5.454 -5.735 19.971 1.00 26.30 A
ATOM	51	CG	GLN	121	-5.128 -4.943 18.710 1.00 26.84 A
ATOM	52	CD	GLN	121	-4.923 -3.460 18.949 1.00 27.26 A
ATOM	53	OE1	GLN	121	-5.822 -2.668 18.709 1.00 28.66 A
ATOM	54	NE2	GLN	121	-3.717 -3.100 19.341 1.00 33.90 A
ATOM		HE21		121	2.883 -3.614 19.564 1.00 15.00 A
ATOM	56	HE22		121	-3.442 -2.138 19.204 1.00 15.00 A
ATOM	57	::===	GLN	121	-8.065 -5.218 19.903 1.00 26.33 A
ATOM	58	Č	GLN	121	-8.905 -6.097 19.834 1.00 21.41 A
ATOM	59	N	ELE	122	-8.288 -4.051 19.272 1.00 21.21 A

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FIGURE 1B

ATOM	60	H	321	122	-7.600	-3.320	19.337		À
			ΞΞΞ	122					
ATOM	51	CA			-9.383	-3.952	18.295		Ä
ATOM	62	CЗ	ILE	122	-10.238	-2.629	18.396	1.00 20.17	À
ATOM	63	CG2	ILE	122	-11.275	-2.428	17.272	1.00 21.61	à
	64	CG:	ILE	122	-11.076	-2.744	19.668	1.00 24.13	Ä
ATOM									
ATOM	65	CD1	ILE	122	-11.751	-1.440	20.073	1.00 23.04	÷
ATOM	66	C	ILE	122	-8.833	-4.108	16.895	1.00 18.96	A
ATOM	67	0	ILE	122	-8.135	-3.243	16.379	1.00 17.93	۸
					-9.159				
ATOM	68	N	ALA	123		-5.240	16.283	1.00 14.72	Ä
ATOM	69	H	ALA	123	-9.599	-5.978	16.805	1.00 15.00	À
ATOM	70	CA	ALA	123	-8.656	-5.401	14.917	1.00 14.29	A
ATOM	71	CB	ALA	123	-7.176	-5.868	14.903	1.00 12.83	Ä
ATOM	72	C	ALA	123	-9.483	-6.315	13.985	1.00 15.66	A
ATOM	73	0	ALA	123	-10.170	-7.261	14.323	1.00 13.58	À
ATOM	74	N	ALA	124	-9.388	-6.009	12.724	1.00 13.45	Ä
ATOM	75	н	ALA	124	-8.894	-5.185	12.456	1.00 15.00	A
									Ä
ATOM	76	CA	ALA	124	-10.087	-6.920	11.836	1.00 14.55	
ATOM	77	CB	ALA	124	-11.486	-6.368	11.446	1.00 11.37	A
ATOM	78	C	ALA	124	-9.271	-7.123	10.563	1.00 13.54	A
ATOM	79	Ō	ALA	124	-8.501	-6.274	10.129	1.00 16.29	A
MOTA	80	N	HIS	125	-9.544	-8.248	9.937	1.00 11.49	A
ATOM	81	Н	HIS	125	-10.100	-B.900	10.426	1.00 15.00	A
ATOM	82	CA	HIS	125	-9.100	-8.524	8.590	1.00 11.51	A
ATOM	83	CB	HIS	125	-7.605	-8.908	B.614	1.00 11.43	A
ATOM	84	CG	HIS	125	-7.119	-9.116	7.205	1.00 7.41	A
ATOM	85	ND1	HIS	125	-6.750	-B.130	6.421	1.00 6.60	A
ATOM	86	HD1	HIS	125	-6.708	-7.168	6.621	1.00 15.00	A
	87		HIS	125	-7.075		6.456	1.00 12.36	A
ATOM									
ATOM	88		HIS	125	-6.670	-9.971	5.234	1.00 6.20	A
ATOM	89	CEl	HIS	125	-6.462	-8.646	5.211	1.00 4.48	A
ATOM	90	С	HIS	125	-10.024	-9.570	7.931	1.00 12.63	A
	91	ō	HIS	125		-10.650	8.383	1.00 13.14	A
ATOM									
MOTA	92	N	VAL	126	-10.550	-9.129	6.806	1.00 15.65	A
ATOM	93	Н	VAL	126	-10.169	-8.286	6.428	1.00 15.00	A
ATOM	94	CA	VAL	126	-11.743	-9.717	6.201	1.00 14.38	A
ATOM	95	CB	VAL	126	-12.877	-8.808	6.675	1.00 13.37	A
ATOM	96	CG1		126	-13.794	-9.722	7.379	1.00 12.60	A
MOTA	97	CG2	VAL	126	-13.449	-7.663	5.814	1.00 9.61	A
ATOM	98	C	VAL	126	-11.502	-9.971	4.685	1.00 16.03	A
ATOM	99	ō	VAL	126	-10.684	-9.297	4.074	1.00 16.42	A
					_		_		
ATOM	100	N	ILE	127	-12.118		4.136	1.00 15.99	A
ATOM	101	H	ILE	127	-12.807	-11.481	4.691	1.00 15.00	A
ATOM	102	CA	ILE	127	-11.651	-11.532	2.831	1.00 14.86	A
ATOM	103	CB	ILE	127	-11.414	-13.051	3.002	1.00 17.56	A
		_			-11.716			1.00 17.17	Ä
MCTA	104	CG2	ILE	127		-13.910	1.765		
ATOM	105	CG:	ILE	127	-9.972	-13.316	3.399	1.00 16.47	A
ATOM	106	CD1	ILE	127	-9.705	-12.992	4.864	1.00 19.64	A
ATOM	107	C	ILE	127	-12.691	-11.269	1.765	1.00 18.96	Α
						-11.391	2.016	1.00 20.01	A
ATOM	108	0	ILE	127					
ATOM	109	N	SER	125		-10.882	0.581	1.00 17.54	A
ATOM	110	H	SER	128	-11.232	-10.871	0.382	1.00 15.00	A
ATOM	111	CA	SER	128		-10.667	-0.437	1.00 15.55	A
				128	-12.664	-10.130			
ATOM	112	CB	SER				-1.706	1.00 18.16	A
MCTA	::3	OG	SER	128		-11.207	-2.574	1.00 19.90	A
ATOM	114	HG	SER	128	-11.832	-11.931	-2.029	1.00 15.00	A
ATOM	115	c	SER	129		-11.761	-0.792	1.00 13.62	A
	116		SER	128		-12.960	-0.832	1.00 8.98	Ä
ATOM	0	C							ň
MOTA	117	ν.	GLU	129		-11.246	-1.027	1.00 13.36	A
ATOM	118	H	SLU	129	-15.661	-10.257	-0.937	1.00 15.00	A
ATOM	119	ΞÀ	SLU	129		-12.024	-1.840	1.00 17.20	A
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FIGURE 1C

MCTA	:2:	CB	SLU	129	-17.052	-13.117	-1.021	1.00 00 55	A
	12:	cs	SLU	129		-12.694		1.00 17.92	Ä
ATOM								1.00 17.92	
ATOM	122	CD	GLU	129	-18.781		C.375		Ä
ATOM	123	SE:	GLÜ	129	-19.997		0.369	1.00 30.23	Ä
ATOM	:24	OE2	GLU	129	-18.150	-14.938	0.734	1.00 33.11	Ä
	125	c	GLU	129		-11.409		1.00 17.71	Ä
MOTA									
ATOM	126	0	GLU	129		-10.389		1.00 21.59	Ä
ATOM	127	N	ALA	130	-17.550	-12.145	-3.914	1.00 20.52	Ä
ATOM	128	н	ALA	130	-17.136	-13.057	-3.923	1.00 15.00	A
	129	CA	ALA	130		-11.649		1.00 23.36	À
ATOM									
MOTA	130	CB	ALA	130		-12.633		1.00 19.66	Ä
ATOM	131	С	ALA	130		-11.298	-4.570	1.00 26.86	À
ATOM	132	0	ALA	130	-20.519	-12.022	-3.869	1.00 29.40	À
ATOM	133	N	SER	131	-20.198	-10.086	-4.968	1.00 21.70	A
	134	н	SER	131	-19.515		-5.410	1.00 15.00	A
ATOM									
MOTA	135	CA	SER	131	-21.592		-4.732	1.00 20.04	A
ATOM	136	CB	SER	131	-21.829	-8.266	-4.787	1.00 20.65	À
ATOM	137	OG	SER	131	-23.182	-8.001	-4.435	1.00 15.24	À
ATOM	138	HG	SER	131	-23.329		-4.559	1.00 15.00	A
						-10.501			
ATOM	139	C	SER	131			-5.668	1.00 17.15	A
ATOM	.140	0	SER	131		-10.853	-6.786	1.00 14.30	A
ATOM	141	N	SER	132	-23.756	-10.731	-5.187	1.00 20.15	A
ATOM	142	H	SER	132	-23.967	-10.586	-4.209	1.00 15.00	А
ATOM	143	CA	SER	132		-11.250	-6.218	1.00 21.62	A
ATOM	144	CB	SER	132		-12.616	-5.893	1.00 16.00	A,
ATOM	145	OG	SER	132	-26.203	-12.324	-4.894	1.00 23.84	A
ATOM	146	HG	SER	132	-26.016	-12.944	-4.179	1.00 15.00	A
MOTA	147	С	SER	132	-25.727	-10.268	-6.671	1.00 20.07	Α
			SER	132		-10.544	-7.547	1.00 20.27	A
ATOM	148	0						-	
MOTA	149	N	LYS	133	-25.606	-9.063	-6.118	1.00 21.87	A
ATOM	150	H	LYS	133	-24.904	-8.969	-5.397	1.00 15.00	A
ATOM	151	CA	LYS	133	-26.406	-7.916	-6.517	1.00 19.23	A
ATOM	152	CB	LYS	133	-27.024	-7.309	-5.256	1.00 23.08	A
			LYS	133	-27.684	-8.364		1.00 21.07	A
ATOM	153	CG					-4.354		
ATOM	154	CD	LYS	133	-29.174	-B.110	-4.320	1.00 27.36	A
ATOM	155	CE	LYS	133	-29.939	-7.884	-5.670	1.00 30.56	Α
ATOM	156	N2	LYS	133	-31.323	-7.515	-5.345	1.00 21.56	А
ATCM	157	HZ1	LYS	133	-31.862	-7.351	-6.218	1.00 15.00	A
ATOM	158		LYS	133	-31.753	-B.299	-4.811	1.00 15.00	A
ATOM	159		LYS	133	-31.333	-6.654	-4.760	1.00 15.00	A
ATOM	160	С	LYS	133	-25.579	-6.876	-7.194	1.00 20.10	A
ATOM	161	0	LYS	133	-24.378	-6.801	-7.007	1.00 17.94	A
ATOM	162	N	THR	134	-26.260	-6.052	-7.983	1.00 22.95	A
ATOM	163	н	THR	134	- 27.275	-6.130	-8.036	1.00 15.00	A
ATOM	164	CA	THR	134	-25.556	-4.879	-8.561	1.00 27.89	A
ATOM	165	CB	THR	134	-26.498	-4.274	-9.592	1.00 24.59	Ä
ATOM	166	0G1	THR	134	-26.540	-5.037	-10.792	1.00 24.32	A
MOTA	167		THR	134	-26.232	-4.411	-11.456	1.00 15.00	Α
MCTA	168		THR	134	-26.044	-2.897	-9.968	1.00 22.97	A
MOTA	169	C	THR	134	-24.987		-7.559	1.00 32.51	A
ATOM	170	၁	THR	134	-25.658	-3.461	-6.603	1.00 38.43	A
ATOM	171	N	THR	135	-23.717	-3.352	-7.690	1.00 35.98	A
ATOM	172	Ħ	THR	135	-23.292	-3.555	-8.585	1.00 15.00	A
ATOM	- 3	Ċλ	THR	135	-22.964	-3.469		1.00 36.02	A
	:;	SE	THR	135	-21.575	-4.276			Ä
ATCM	- 7	-=					-6.534	1.00 36.01	
ATOM	175	23:	THR	135	-21.645		-7.488	1.00 30.60	A
ATOM	:76	H31	THR	135	-22.255	-6.094	-7.312	1.00 15.00	A
ATOM	:	222	THR	135	-20.866	-4.776	-5.264	1.00 35.55	A
ATOM	- 5	=	THR	135	-22.949		-5.404	1.00 30.25	Ä
ATOM		Ē	THR	135	-23.541	-2.348	-4.331		2
A.U.		J		•	- 63.34.	- 2 . 3 7 6	-4.23.	1.00 28.35	^

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FIGURE 1D

OM	180	::	SER	136	-22.294	-1.146	-5.776	1.33 23.29	A
ATOM		H H	SER	136	-12.828	-0.357	-5.463		Ä
ATOM	151				-20.857				Ä
ATOM	152	CA	SER	136		-1.051	-6.143	1.00 03.04	
ATOM	163	CB	SER	136	-20.560	0.187	-6.965	1.00 21.03	÷
ATOM	184	၁၄	SER	13€	-20.624	1.261	-6.043	1.00 28.21	Ä
MCTA	185	HG	SER	136	-19.815	1.793	-6.008	1.00 15.00	À
ATOM	186	C	SER	13€	-19.553	-1.090	-4.958	1.00 21.77	À
ATOM	187	0	SER	13€	-18.630	-1.096	-5.080	1.00 21.94	À
ATOM	188	N	VAL	137	-20.452	-1.227	-3.752	1.00 24.03	· À
ATOM	189	H	VAL	137	-21.440	-1.063	-3.705	1.00 15.00	À
ATOM	190	CA	VAL	137	-19.699	-1.632	-2.570	1.00 19.65	À
	191	CB	VAL	137	-20.218	-1.010	-1.248	1.00 21.14	 A
ATOM		CG1	VAL	137	-20.419	-1.907	-C.058	1.00 18.16	Ä
ATOM	192		VAL	137	-21.322	-0.026	-1.442	1.00 13.49	
ATOM	193	CG2							Ä
ATOM	194	C	VAL	137	-19.370	-3.116	-2.473	1.00 17.15	À
ATOM	195	0	VAL	137	-20.209	-3.969	-2.593	1.00 16.69	A
ATOM	196	N	LEU	138	-18.077	-3.344	-2.271	1.00 15.84	Α
ATOM	197	H	LEU	138	-17.502	-2.528	-2.246	1.00 15.00	A
ATOM	198	CA	LEU	138	-17.507	-4.667	-1.938	1.00 18.21	A
ATOM	199	CB	LEU	138	-15.962	-4.530	-1.791	1.00 13.60	A
ATOM	200	CG	LEU	138	-15.273	-3.854	-2.998	1.00 16.09	A
ATOM	201		LEU	138	-15.923	-4.379	-4.300	1.00 20.35	A
ATOM	202		LEU	138	-13.710	-3.936	-2.982	1.00 12.34	A
ATOM	203	C	LEU	138	-18.170	-5.480	-0.772	1.00 16.29	A
		Õ	LEU	138	-18.498	-4.986	0.301	1.00 12.97	A
ATOM	204							1.00 13.04	Ä
ATOM	205	N	GLN	139	-18.345	-6.768	-1.035		
ATOM	206	H	GLN	139	-18.052	-7.078	-1.960	1.00 15.00	A
ATOM	207	CA	GLN	139	-18.757	-7.658	0.013	1.00 15.32	À
MOTA	208	CB	GLN	139	-19.847	-8.678	-0.481	1.00 13.99	A
MOTA	209	CG	GLN	139	-21.068	-7.960	-1.113	1.00 20.85	A
ATOM	210	CD	GLN	139	-21.872	-7.022	-0.193	1.00 22.04	A
ATOM	211	OEl	GLN	239	-22.343	-7.439	0.878	1.00 25.45	A
MOTA	212	NE2		139	-21.963	-5.739	-0.618	1.00 17.74	A
ATOM	213	HE21	GLN	139	-22.697	-5.181	-0.206	1.00 15.00	A
ATOM	214	HE22		139	-21.460	-5.326	-1.374	1.00 15.00	A
ATOM	215	C	GLN	139	-17.527	-8.383	0.541	1.00 14.26	Α
ATOM	216	ō	GLN	139	-16.554	-8.640	-0.144	1.00 14.40	A
MCTA	217	N	TRP	140	-17.647	-8.780	1.805	1.00 12.80	A
	218	H	TRP	140	-18.433	-8.447	2.297	1.00 15.00	A
ATOM			TRP					1.00 14.03	Ä
ATOM	219	CA		140	-16.542	-9.500	2.463 3.483		
ATOM	220	CB	TRP	140	-15.813	-8.623		1.00 14.18	A
ATOM	221	CG	TRP	140	-15.467	-7.291	2.823	1.00 8.44	A
ATOM	222	CD2	TRP	140	-14.379	-6.966	1.941	1.00 9.01	À
MOTA	223	CE2	TRP	140	-14.549	-5.625	1.482	1.00 8.40	A
MCTA	224	CE3	TRP	140	-13.215	-7.688	1.581	1.00 10.14	A
MCTA	225	CD:	TRP	140	-16.225	-6.137	2.863	1.00 11.29	A
ATOM	226	NE:	TRP	140	-15.710	-5.150	2.077	1.00 14.27	A
MOTA	227	HE1	TRP	14C	-16.121	-4.268	2.010	1.00 15.00	Α
ATOM	228	CZ2		140	-13.640	-5.009	0.590	1.00 B.16	Α
ATOM	229	CZ3	TRP	140	-12.292	-7.069	0.713	1.00 13.90	A
ATOM	230	CH2	TRP	140	-12.497	-5.749	0.215	1.00 12.11	A
ATOM	231	c	TRP	140		-10.701	3.170	1.00 14.34	Ä
MOTA	232	Š	TRP	140	-18.193	-10.862	3.392	1.00 16.00	Ä
				141	-16.582	-11.528	3.558	1.00 14.80	Ä
ATOM	233	N	<u> </u>	• • •					, A
MCTA	234	Η,	بنند	141	-15.133	-11.377	3.294	1.00 15.00	÷
ATOM	235	<u> </u>	^ -^	141	-16.489	-12.617	4.394	1.00 15.27	Ä
ATOM	236	CB	À-À	141	-16.504	-13.920	3.583	1.00 16.97	Ä
ATOM	237	=	ALA	141		-12.761	5.607	1.00 15.90	A
ATOM	239		نمنند	:4:	-14.453	-12.338	5.550	1.00 14.25	Ä
ATOM	239	N	320	141	-16.069	-13.366	5.688	1.00 19.74	Ä

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FIGURE 1E

17:04	240	H	31C	142	-17.055	-13.574	6.688	1.00 15.00	A
ATOM				142	-15.149		7.731	1.00 05.93	÷
ATOM	241	ΞÀ	<u> </u>				9.117	1.00 25.93	
MOTA	242	= 3	GLU	142		-13.910			Ä
ATOM	243	CG	GLU	142	-15.716	-12.456	5.64	1,00 04.05	Ä
MOTA	244	CD	GLU	142	-15.749	-12.087	10.711	1.00 26.61	Ä
		OE1	GLU	142	-17.908	-11.888	10.361	1.00 34.72	A
ATOM	245					-11.984	11.886	1.00 30.07	Ä
ATOM	246	QE2		142				1.00 33.25	
ATOM	247	C	GLU	142	-14.200		7.193		÷
ATOM	248	0	GLU	142	-13.156	-14.349	6.737	1.00 41.84	Ä
MCTA	249	N	LYS	143	-14.577	-16.080	7.084	1.00 34.17	Ä
	250	H	LYS	143	-15.432	-16.384	7.492	1.00 15.00	A
ATOM						-16.854	5.980	1.00 35.31	Ä
MOTA	251	CA	LYS	143			4.681	1.00 37.64	Ä
ATOM	252	CB	LYS	143	-14.673				
MOTA	253	CG	LYS	143	-14.300		3.531	1.00 47.37	A
ATOM	254	CD	LYS	143	-15.022	-17.284	2.202	1.00 50.37	Ä
ATOM	255	CE	LYS	143	-14.686	-16.047	1.357	1.00 49.23	A
	256	NZ	LYS	143		-16.097	0.221	1.00 51.67	A
MOTA					-15.333	_	-0.534	1.00 15.00	Ä
ATOM	257		LYS	143			-0.177	1.00 15.00	A
ATOM	258	HZ2	LYS	143	-15.680				
ATOM	259	HZ3	LYS	143	-16.564-		0.585	1.00 15.00	Α
ATOM	260	C	LYS	143	-12.330	-16.979	5.637	1.00 32.80	A
ATOM	261	Ó	LYS	143	-11.831	-18.041	5.276	1.00 35.64	A
	262	N	GLY	144	-11.522		5.637	1.00 28.26	A
MOTA					-11.718		5.910	1.00 15.00	A
ATOM	263	H	GLY	144			5.194	1.00 32.94	A
ATOM	264	CA	GLY	144	-10.243				
ATOM	265	C	GLY	144		-16.862	6.180	1.00 29.93	A
ATOM	266	0	GLY	144		-17.454	7.205	1.00 24.67	A
ATOM	267	N	TYR	145	-0.069	-16.270	5.815	1.00 26.37	À
	268	н	TYR	145	-8.160	-15.729	4.966	1.00 15.00	A
ATOM			TYR	145		-16.002	6.777	1.00 27.61	A
ATOM	269	CA				-15.877	5.947	1.00 37.54	A
ATOM	270	CB	TYR	145				1.00 50.95	Ä
MCTA	271	CG	TYR	145		-15.774	4.456		
ATOM	272	CD1	TYR	145		-14.633	3.706	1.00 53.22	Ä
ATOM	273	CE1	TYR	145	-6.313	-14.377	2.468	1.00 60.28	A
ATOM	274	CD2	TYR	145	-6.591	-16.847	3.791	1.00 53.11	A
MOTA	275	CE2		145	-7.207	-16.699	2.551	1.00 56.30	~
			TYR	145		-15.430	1.873	1.00 61.12	A
ATOM	276	CZ				-15.119	0.665	1.00 62.63	A
MCTA	277	OH	TYR	145				1.00 15.00	A
MOTA	27B	HH	TYR	145		-15.686	0.401		
ATOM	279	C	TYR	145		-14.762	7.620	1.00 22.41	Ä
ATOM	280	C	TYR	145	-7.000	-13.677	7.650	1.00 22.68	A
ATOM	281	N	TYR	146	-8.731	-14.884	B.196	1.00 20.39	A
ATOM	282	н	TYR	146	-8.935	-15.824	8.509	1.00 15.00	A
		CA	TYR	:46		-13.700	8.725	1.00 20.40	A
MOTA	283			146		-13.673	B.306	1.00 22.53	A
ATOM	284	CB	TYR				9.286	1.00 23.02	Ä
ATOM	285	CG	TYR	146	•11.715	-14.460			
ATOM	296		TYR	146	•11.635	-15.873	9.236	1.00 26.99	A
ATOM	287	CE:	TYR	146		-16.623	10.239	1.00 25.44	A
ATOM	288	CD2		146	-12.477	-13.766	10.236	1.00 23.45	Ä
	289			146		-14.520	11.205	1.00 26.81	A
MOTA				146		-15.937	11.204	1.00 27.40	A
ATOM	290	CZ	TYR			-16.689	12.170	1.00 31.91	A
ATOM	291	OH	TYR	146	3.04/	10.007			Ä
ATOM	292	HH	TYP	146		-17.080	12.676	1.00 15.00	•
MOTA	293	0	TYR	:4€		-13.419	10.219	1.00 18.79	÷
ATOM	294	2	TYR	146	- 3 . 904	-14.232	11.012	1.00 16.13	Ä
MCTA	295	::	THE	147		-12.169	10.556	1.00 17.54	A
	296	H	THR	147		-11.607	9.830	1.00 15.00	Ä
ATOM				: 47		-11.764	11.948	1.00 14.06	Ä
ATOM	297	ÇA	THR				12.182	1.00 13.65	Ä
MOTA	295	23	THR	147		-10.875			Ä
ATOM	299	23	THR	:47	-5.512	-11.505	11.856	1.00 12.56	~

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FIGURE 1F

ATOM	300 H3	: THR	147	- é . 934	-11.995	10.980		Ä
MCTA	301 03	I THR	147		-10.236	13.554	1:3: 7:22	2
ATOM	302 0	THR	147		-10.925	12.253		
ATOM	303 C	THR	147		-10.074	11.496	1,00 16.39	÷
ATOM	304 N	MET	145		-11.139		1.00 16.39	è
MOTA	305 H	MET	148		-11.988	13.412	1.00 20.67	Ä
ATOM	306 CA			12.836	-11.988	13.828	1.00 15.00	Ä
			148		-10.311	14.110	1.00 19.71	À
ATOM	367 CB		148		-10.702	13.705	1.00 17.89	A
ATOM	308 CG		148	-14.541	-9.580	14.019	1.00 13.53	A
MOTA	309 SD		148	-14.492	-8.149	12.952	1.00 14.69	A
ATOM	310 CE	MET	148	-14.566	-8.928	11.333	1.00 10.10	A
ATOM	311 C	MET	148		-10.282	15.639	1.00 21.49	Ä
ATOM	312 0	MET	148	-12.594	-10.905	16.436	1.00 22.98	A
ATOM	313 N	SER	149	-10.955	-9.412	16.055	1.00 20.58	Ä
ATOM	314 H	SER	149	-10.516	-8.786	15.406	1.00 15.00	Ä
ATOM	315 CA	SER	149	-10.388	-9.698	17.419	1.00 19.11	 A
ATOM	316 CB	SER	149	-9.174	-8.860	17.792	1.00 12.17	Ä
ATOM	317 OG	SER	149	-9.540	-7.513	17.975	1.00 14.10	Â
ATOM	318 HG	SER	149	-9.571	-7.487	18.934	1.00 15.00	Ä
ATOM	319 C	SER	149	-11.203	-9.844	18.727	1.00 22.19	
ATOM	320 0	SER	149	-10.728		19.772	1.00 22.95	A
ATOM	321 N	ASN	150	-12.456	-9.322	18.631		A
ATOM	322 H	ASN	150	-12.782	-9.247		1.00 22.71	À
ATOM	323 CA	ASN	150	-13.361		17.688	1.00 15.00	A
ATOM	324 CB	ASN	150		-9.236	19.764	1.00 20.32	A
ATOM	325 CG	ASN	150	-12.734	-8.446	20.955	1.00 21.56	A
ATOM		ASN		-12.343	-6.962	20.706	1.00 20.71	A
ATOM		ASN ASN	150	-13.059	-6.187	20.119	1.00 17.81	A
			150	-11.222	-6.485	21.271	1.00 23.86	A
ATOM		ASN	150	-11.035	-5.521	21.092	1.00 15.00	Α
ATOM		2 ASN	150	-10.670	-7.109	21.821	1.00 15.00	A
ATOM	330 C	ASN	150	-14.644	-8.657	19.256,	1.00 20.60	A
ATOM	331 0	ASN	150	-14.718	-8.130	18.148	1.00 20.56	A
MCTA	332 N	ASN	151	-15.637	-8.713	20.149	1.00 23.49	A
ATOM	333 H	asn	151	-15.455	-9.124	21.038	1.00 15.00	À
ATOM	334 CA	asn	151	-16.974	-8.080	19.823	1.00 24.71	A
ATOM	335 CB	ASN	151	-18.130	-8.645	20.712	1.00 28.30	A
ATOM	336 CG	asn	151	-17.959	-8.271	22.173	1.00 33.23	A
ATOM		ASN	151	-17.075	-7.562	22.606	1.00 39.79	A
ATOM		ASN	151	-18.782	-8.838	23.011	1.00 38.32	A
MOTA		. ASN	151	-18.553	-8.524	23.928	1.00 15.00	À
MCTA	340 HD22	ASN	151	-19.495	-9.465	22.733	1.00 15.00	Ä
ATOM	341 C	ASN	151	-17.172	-6.531	19.645	1.00 22.53	Ä
MOTA	3 4 2 O	ASN	151	-18.254	-6.048	19.374	1.00 21.32	Ä
ATCM	343 N	LEU	152	-16.066	-5.762	19.859	1.00 23.00	Ä
ATOM	344 H	LEU	152	-15.247	-6.289	20.070	1.00 15.00	Ä
ATOM	345 CA	LEU	152	-15.924	-4.335	19.525	1.00 18.87	Ä
ATOM	346 CB	LEU	152	-14.830	-3.700	20.325	1.00 21.77	
MCTA	347 CG	LEU	152	-14.981	-3.999	21.806	1.00 24.80	Ä
MCTA	348 CD1		152	-16.390	-3.645	22.316	1.00 22.82	Ä
MCTA	349 CD2	LEU	152	-13.847	-3.256	22.556	1.00 23.56	
ATOM	350 C	LEU	152	-15.565	-3.993	18.094		À
ATOM	351 0	LEU	152	-15.590	-2.840		1.00 17.34	A
ATOM	352 N	VAL	153	-15.267		17.708	1.00 13.39	Ą
ATOM	353 H	VAL	153		-5.054	17.309	1.00 18.65	A
MCTA	354 CA	VAL	153	-15.156	-5.962	17.716	1.00 15.00	À
ATOM				-15.439	-4.910	15.849	1.00 16.81	À
ATOM	366 CG:	VAL	153	-14.138	-5.021	14.980	1.00 15.33	À
ATOM			153	-12.908	-5.718	15.562	1.00 21.22	Ä
7 TON			153	-13.775	-3.757	14.287	1.00 16.95	Ä
ATOM	358 0	VAL	153	-16.405	-5.964	15.30:	1.00 13.48	A
ATOM	359 C	VAL	153	-16.363	-7.:16	15.647	1.00 13.06	Ä

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FIGURE 1G

ATOM	340 N	THE	154	-17.207	-5.546	14.358		:	•
ATOM	361 H	THE	154	-17.313	-4.568	14.215			À
								.E.::	À
ATOM	362 CA		154	-17.903	-6.600	13.615	1.00	. .	Ä
MCTA	363 CB	THR	154	-19.366	-6.747	14.157	1.75	, S ', E :	A
MCTA	364 CG	I THR	. 154	-19.995	-5.459	14.205	1.00	4 ::	Ä
ATOM	3 6 5 HG		154	-20.577	-5.508	14.949			
							1.0C 1	5.00	Ä
ATOM	366 CG		154	-19.502	-7.288	15.571		1.62	Ä
MOTA	367 C	THR	154	-17.997	-6.252	12.107	1.00 1	5.12	
MOTA	368 O	THR	154	-17.992	-5.110	11.605	1.00 1		*
ATOM	369 N	LEU	155	-18.101	-7.324	11.357	1.00 1		
		LEU							À
ATOM			155	-18.056	-8.202	11.791	1.00 1		A
ATOM	371 CA		155	-18.514	-7.198	9.967	1.00 1	7.10	A
MOTA	372 CB	LEU	155	-17.829	-8.353	9.204	1.00 2	C.04	Ä
MOTA	373 CG	LEU	155	-17.524	-8.428	7.692	1.00 2	0.8	Ä
ATOM	374 CD	1 LEU	155	-17.822	-7.159	6.908	1.00 1	_	
ATOM		2 LEU	155	-17.912	-9.810				÷
						7.139	1.00 1		Ä
ATOM	376 C	LEU	155	-20.055	-7.187	9.904	1.00 2		Ä
ATOM	377 0	LEU	155	-20.712	-8.163	10.217	1.00 1	8.01	A
ATOM	378 N	GLU	15€	-20.593	-5. 995	9.561	1.00 1	9.51	À
ATOM	379 H	GLU	156	-19.959	-5.230	9.440	1.00 1		Ä
ATOM	380 CA		156	-22.036					
					-5.888	9.413	1.00 2		A
ATOM	381 CB	GLU	156	-22.641	-4.631	10.033	1.00 1	8.95	A
ATOM	382 CG	GLU	156	-22.098	-4.412	11.436	1.00 2	7.68	A
ATOM	383 CD	GLU	156	-22.721	-5.194	12.587	1.00 3	1.62	A
ATOM	384 CE	1 GLU	156	23.347	-6.248	12.367	1.00 3		A
ATOM		2 GLU	156	-22.532	-4.721	13.724	1.00 3		
ATOM	386 C	GLU	156						A
				-22.457	-5.966	7.964	1.00 2		A
ATOM	387 O	GLU	156	-21.958	-5.298	7.077	1.00 2	2.70	À
ATOM	388 N	ASN	157	-23.437	-6.808	7.696	1.00 3	0.92	Α
ATOM	389 H	ASN	157	-23.594	-7.590	8.300	1.00 1	5.00	A
ATOM	390 CA	ASN	157	-23.804	-6.620	6.300	1.00 3		Ä
ATOM	391 CB	ASN	157	-23.856					
A TOM					-7.970	5.614	1.00 3		Ą
MCTA	392 CG	ASN	157	-23.669	-7.693	4.168	1.00 2	7.70	A
ATOM		1 ASN	157	-23.397	-6.593	3.810	1.00 2	5.89	A
MCTA	394 ND:	2 ASN	157	-23.893	-B.640	3.275	1.00 4	1.69	A
MOTA	395 HD2:	i ASN	157	-24.069	-9.603	3.467	1.00 1	5 . 00	A
ATOM	396 HD2:	2 ASN	157	-23.745	-8.295	2.340	1.00 1		Ä
ATOM	397 C	ASN	157	-24.988	-5.658	6.118			
ATOM							1.00 3		Ą
		ASN	157	-26.107	-5.949	6.499	1.00 3		A
MOTA	399 N	GLY	158	-24.746	-4.443	5.560	1.00 4	0.03	À
ATOM	400 H	GLY	15ê	-25.601	-3.952	5.429	1.00 1	5.00	À
ATOM	401 CA	GLY	158	-23.422	-3.887	5.121	1.00 3	R 11	A
ATOM	492 C	GLY	158	-23.062	-3.720	3.617	1.00 3		Ä
ATOM	403 0	GLY	158	-23.890	-3.108				
ATOM						2.950	1.00 4		A
	404 N	LYS	155	-11.867	-4.220	3.135	1.00 3	2.75	À
atom	405 H	LYS	159	- 21 . 904	-4.134	2.130	1.00 1	5.00	Ä
MOTA	406 CA	LYS	159	-20.829	-4.928	3.962	1.00 2	7.83	A
ATOM	407 CB	LYS	159	-20.317	-6.122	3.217	1.00 2		A
ATOM	408 CG	LYS	159	-19.734	-7.168	4.069	1.00 2		
MCTA	409 CD	LYS	159						À
				-20.533	-B.426	4.192	1.00 2		A
MCTA	410 CE	LYS	159		-9.191	2.869	1.00 4		A
MOTA	411 NZ	LYS	159		-10.663	2.986	1.00 4	0.88	A
ATOM	412 HZ		159		-11.087	2.035	1.00 1		À
MCTA	413 HZ	1115	159		-11.087	3.600	1.00 1		Ä
ATOM	414 HZ		: 5 5		-10.848				•
ATOM	415 7					3.389	1.00 1		÷
A . C.	415 0	LYS	159	-19.688	-4.065	4.463	1.00 2		Ä
ATOM	416 0	LYS	159	-19.023	-3.369	3.696	1.00 2		÷
ATOM	417 N	31.N	160	-19.683	-3.990	5.807	1.00 1		Ä
ATOM	418 H	31.:	160	-20.211	-4.674	6.3.9	1.00 1		Ä
ATOM	419 CA		160	-19.922	-2.929	6.464	1.00 1		
						U . T U 7	1.00	J. C7	~

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FIGURE 1H

MOTA	420	CB	3LN	160	-19.778	-1.694	6.611	1.00 16.79	Ä
				160	-20.881	-1.896	7.633		
ATOM	421	CS	GLN					1.00 18.34	÷
MCTA	422	CD	GLN	160	-22.133	-1.166	7.193	1.00 23.97	Ä
ATOM	423	OE1	GLN	160	-23.086	-0.970	7.893	1.00 31.18	A
	424	NE2	GLN	160	-22.257	-C.771	5.948	1.00 28.16	Ä
ATOM									
ATOM	425	HE21	GLN	160	-23.194	-0.420	5.928	1.00 15.00	À
MCTA	425	HE22	GLN	16C	-21.624	-0.780	5.186	1.00 15.00	Ä
			GLN	160	-18.313	-3.309	7.777		
MOTA	427	<u> </u>						1.00 12.67	À
ATOM	428	0	GLN	160	-18.838	-4.151	8.498	1.00 14.78	À
ATOM	429	N	LEU	161	-17.187	-2.637	8.085	1.00 11.22	À
	430	н	LEU	161	-16.767	-2.124	7.340	1.00 15.00	Ä
ATOM									
ATOM	431	CA	LEU	161	-16.583	-2.870	9.405	1.00 9.71	À
ATOM	432	CB	LEU	161	-15.052	-2.939	9.390	1.00 4.67	À
ATOM	433	CG	LEU	161	-14.438	-4.060	8.559	1.00 7.30	À
MOTA	434		LEU	161	-14.511	-5.447	9.207	1.00 10.80	A
ATOM	435	CD2	LEU	161	-12.964	-3.794	8.389	1.00 5.48	Α
ATOM	436	С	LEU	161	-17.082	-1.836	10.412	1.00 10.17	Α
		ŏ	LEU	161	-16.826			1.00 13.36	Ä
ATOM	437					-0.657	10.341		
ATOM	43B	N	THR	162	-17.848	-2.338	11.375	1.00 16.94	A
ATOM	439	Н	THR	162	-18.153	-3.279	11.251	1.00 15.00	Α
ATOM	440	CA	THR	162	-18.317	-1.480	12.493	1.00 16.14	A
ATOM	441	CB	THR	162	-19.807	-1.769	12.640	1.00 13.33	A
ATOM	442	0G1	THR	162	-20.339	-1.707	11.308	1.00 16.73	A
ATOM	443	HG1	THR	162	-21.211	-1.254	11.343	1.00 15.00	Α
MOTA	444	CG2		162	-20.553	-0.832	13.562	1.00 15.01	A
MOTA	445	C	THR	162	-17.531	-1.547	13.842	1.00 13.28	A
ATOM	446	0	THR	162	-17.358	-2.587	14.449	1.00 20.21	A
ATOM	447	N	VAL	163	-16.994	-0.437	14.282	1.00 14.22	A
									_
ATOM	448	н	VAL	163	-16.859	0.243	13.567	1.00 15.00	A
ATOM	449	CA	VAL	163	-16.326	-0.358	15.586	1.00 15.72	Α
ATOM	450	CB	VAL	163	-15.038	0.426	15.428	1.00 11.82	Α
ATOM	451	CG1	VAL	163	-15.191	1.944	15.368	1.00 9.87	A
ATOM	452			163	-14.229	-0.124	14.245	1.00 18.88	A
ATOM	453	С	VAL	163	-17.193	0.283	16.706	1.00 17.93	A
MOTA	454	0	VAL	163	-18.001	1.180	16.453	1.00 20.25	A
ATOM	455	N	LYS	164	-17.037	-0.232	17.925	1.00 15.44	A
ATOM	456	H	LYS	164	-16.254	-0.858	18.020	1.00 15.00	A
ATOM	457	CA	LYS	164	-17.856	0.138	19.109	1.00 17.33	A
MCTA	458	CB	LYS	164	-18.351	-1.150	19.807	1.00 19.58	A
ATOM	459	ĊĠ	LYS	164	-19.214	-1.885	18.759	1.00 23.56	A
ATOM	460	CD	LY5	164	-19.417	-3.410	18.851	1.00 28.85	A
ATOM	461	CE	LYS	164	-20.039	-4.047	17.554	1.00 33.81	A
ATOM	462	NZ	LYS	164	-19.428	-3.681	16.227	1.00 18.98	A
ATOM	463		LYS	164	-19.195	-2.667	16.222	1.00 15.00	A
ATOM	464		LYS	164	-10.552	-4.223	16.092	1.00 15.00	A
MCTA	465	HZ3	1 YS	164	·20.084	-3.888	15.445	1.00 15.00	A
MOTA	466	C	LYS	164	-17.193	1.099	20.056	1.00 15.14	A
	467	_		164	-17.712	1.588			
MOTA		0	LYS				21.048	1.00 17.72	A
MOTA	468	N	ARG	165	-15.992	1.428	19.621	1.00 17.49	A
ATOM	469	H	ARG	165	-15.550	0.838	18.932	1.00 15.00	Α
ATOM	470	CA	ARG	165	-15.184	2.415	20.325	1.00 20.18	A
MOTA	471	CB	ARG	165	-13.985	1.806	21.049	1.00 24.65	A
MOTA	472	CG	ARG	165	-14.363	0.833	22.126	1.00 29.54	A
ATOM	473	CD	ARG	165	-13.274	1.077	23.145	1.00 38.82	A
ATOM	474	NE	ARG	165	-13.719	1.998	24.186	1.00 43.41	A
		HE.							
ATOM	475		ARS	165	-14.331	1.671	24.908	1.00 15.00	. A
ATOM	47é	CZ	ARG	165	-13.190	3.250	24.362	1.00 44.06	A
ATOM	477	NH1	ARG	165	-13.406	3.765	25.562	1.00 41.25	A
ATOM		HHII		165	-13.054	4.683	25.763	1.00 15.00	Ä
		HHIZ							
ATOM	4 / 7	nn	MK O	165	-13.919	3.249	26.250	1.00 15.00	Ä

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FIGURE 1I

ATOM	4 B C	NH2	ARG	165	-12.485	3.946	23.425	1.00 31.65	A
MCTA	481	HH21		165	-12.133	4.860	23.623	1.00 15.00	
ATOM	482	HH22	ARG	165	-12.322	3.527	22.530	1.00 15.00	
ATOM	483	Ξ	ARG	165	-14.608	3.554	19.510	1.00 17.70	Ä
ATOM	484	С	ARG	165	-14.018	3.450	18.441	1.00 18.26	Ä
ATOM	485	N	GLN	166	-14.763	4.687	20.151	1.00 17.43	
ATOM	486	Н	GLN	166	-15.263	4.614	21.007	1.00 15.00	
ATOM	487	CA	GLN	166	-14.138	5.911	19.698	1.00 19.00	
MOTA	489	CB	GLN	155	-14.613	7.021	20.610	1.00 23.79	
ATOM	489	CG	GLN	166	-14.067	8.409	20.386	1.00 34.06	À
ATOM	490	CD	GLN	166	-15.178	9.399	20.659	1.00 45.91	À
ATOM	491		GLN	166	-15.102	10.492	20.135	1.00 53.64	
ATOM	492		GLN	166	-16.202	9.046	21.418	1.00 44.10	
ATOM		HE21		166	-16.906	9.765	21.443	1.00 15.00	A
ATOM		HE22		166	-16.577	8.287	21.935	1.00 15.00	À
ATOM	495	С	GLN	166	-12.649	5.881	19.644	1.00 17.48	A
ATOM	496	0	GLN	166	-12.029	5.378	20.561	1.00 18.13	A
ATOM	497	N	GLY	167	-12.160	6.478	18.565	1.00 14.83	A
ATOM	49B	H	GLY	167	-12.750	6.836	17.850	1.00 15.00	A
ATOM	499	CA	GLY	167	-10.728	6.711	18.557	1.00 16.28	A
MOTA	500	C	GLY	167	-10.044	6.685	17.204	1.00 16.48	A
ATOM	501	0	GLY	167	-10.674	6.601	16.162	1.00 19.19	A
ATOM	502	N	LEU	168	-8.720	6.735	17.209	1.00 17.06	A
ATOM	503	H	LEU	168	-8.311	6.890	18.120	1.00 15.00	
ATOM	504	CA	LEU	168	-7.925	6.625	15.992	1.00 16.60	
ATOM	505	CB	LEU	168	-6.600	7.343	16.289	1.00 21.87	
ATOM	506	CG	LEU	168	-6.247	8.745	15.716	1.00 22.69	A
ATOM ATOM	507 508		LEU	168	-5.119	9.410	16.539	1.00 21.20	A
ATOM	509	CD2	LEU	168 168	-7.436 -7.686	9.617	15.361 15.604	1.00 18.38	A
ATOM	510	Ö	LEU	168	-7.282	5.136 4.278		1.00 14.84	A
ATOM	511	N	TYR	169	-7.943	4.278	16.392 14.300	1.00 15.89	A
MCTA	512	н	TYR	169	-8.313	5.659	13.807	1.00 10.57	Ä
MOTA	513	ĊA	TYR	169	-7.683	3.572	13.656	1.00 15.00 1.00 5.27	A A
MOTA	514	CB	TYR	169	-8.989	3.014	13.230	1.00 5.83	À
ATOM	515	ČĞ	TYR	169	-9.857	2.620	14.423	1.00 6.94	À
ATOM	516	CD1	TYR	169	-10.524	3.598	15.168	1.00 7.40	Ä
ATOM	517	CE1	TYR	169	-11.390	3.193	16.218	1.00 7.77	Ä
MOTA	518	CD2	TYR	169	-10.016	1.255	14.744	1.00 8.89	Ä
ATOM	519	CE2	TYR	169	-10.850	0.841	15.804	1.00 9.40	
MOTA	520	CZ	TYR	169	-11.563	1.827	16.534	1.00 10.39	Ä
MOTA	521	OH	TYR	169	-12.443	1.410	17.534	1.00 7.99	A
MOTA	522	HH	TYR	169	-13.009	2.117	17.800	1.00 15.00	A
MOTA	523	C	TYR	169	-6.810	3.642	12.390	1.00 6.72	A
ATOM	524	0	TYR	169	-6.917	4.498	11.557	1.00 9.12	A
ATOM	525	N	TYR	170	-5.899	2.722	12.228	1.00 9.53	A
ATOM	526	Н	TYR	170	-5.806	2.081	12.986	1.00 15.00	A
MOTA	527	CA	TYR	170	-5.313	2.511	10.899	1.00 10.01	A
ATOM	528	CB	TYR	170	-3.967	1.797	11.044	1.00 7.46	À
MOTA	529	CG	TYR	170	-3.259	1.636	9.679	1.00 13.45	- A
ATOM	530	CD:	TYR	170	-2.680	2.766	9.052	1.00 12.66	A
ATOM	531	CEL	TYR	170	-2.213	2.658	7.738	1.00 10.18	A
MCTA	532	CD2	TYR	170	-3.304	0.385	9.057	1.00 10.90	
MOTA	533	CE2	TYR	170	-2.891	0.303	7.730	1.00 8.68	
ATOM	534	CZ	TYR	176	-2.331	1.419	7.124	1.00 9.97	
ATOM	535	OH	TYR	170	-1.774	1.286	5.859	1.00 17.50	
ATOM	536	HH	TYR	170	-1.886	0.404	5.514	1.00 15.00	
MOTA	537	Ξ	TYR	170	-6.279	1.€10	10.073	1.00 10.40	
MOTA	53B	0	TYR	170	-6.679	0.500	10.421	1.00 12.52	
MCTA	539	N	ILE	:71	-5.704	2.174	8.962	1.00 12 16	<u>.</u>

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FIGURE 1J

MCTA	54 C	Ξ	ΞLΕ	171	-6.475	3.135	8.808	• • • • • • • • • • • • • • • • • • • •	
	541	CA	ΞΞΞ		-7.505	1.430	8.136	1.00 9.37	Ä
ATOM								7.3	
MCTA	542	C3	ILΕ	171	-9.070	1.990	8.317	1.11 11.11	A
ATOM	543	CG2	ILE	:7:	-9.326	3.501	8.677	1.00 17.00	Ä
ATOM	544	CG1	ILE	171	-10.046	1.564	7.214	1.00 13.33	Ä
ATOM	545	CDI	ILE	171	-10.647	0.250	7.619	1.00 17.53	Ä
ATOM	546	C	ILE	171	-7.074	1.234	6.694	1.00 9.34	Ä
ATOM	547	0	ILE	171	-6.453	2.088	6.082	1.00 6.96	A
ATOM	548	N	TYR	172	-7.286	0.005	6.216	1.00 11.07	Ä
ATOM	549	H	TYR	172	-7.809	-0.624	6.786	1.00 15.00	Ä
		-	TYR	172	-6.708	-0.378	4.922	1.00 15.60	Ä
ATOM	550	CA							
ATOM	551	CB	TYR	172	-5.332	-1.082	5.037	1.00 14.32	Ä
ATOM	552	CG	TYR	172	-5.389	-2.397	5.796	1.00 9.21	Ä
ATOM	553	CD1	TYR	172	-5.342	-2.402	7.216	1.00 12.52	À
ATOM	554	CE1	TYR	172	-5.607	-3.620	7.901	1.00 10.89	À
ATOM	555	CD2	TYR	172	-5.565	-3.586	5.050	1.00 12.66	Ä
ATOM	556	CE2	TYR	172	-5.829	-4.800	5.740	1.00 15.83	A
ATOM	557	CZ	TYR	172	-5.822	-4.808	7.164	1.00 11.94	A
ATOM	558	OH	TYR	172	-5.995	-6.002	7.820	1.00 12.17	Α
ATOM	559	HH	TYR	172	-6.433	-5.843	8.657	1.00 15.00	A
ATOM	560	C	TYR	172	-7.605	-1.276	4.106	1.00 16.85	A
	561	Ö	TYR	172	-8.346	-2.057	4.692	1.00 14.06	A
ATOM									
MOTA	562	N	ALA	173	-7.448	-1.141	2.776	1.00 16.29	λ
ATOM	563	Н	ALA	173	-6.751	-0.490	2.503	1.00 15.00	A
ATOM	564	CA	ALA	173	-7.940	-2.152	1.836	1.00 15.11	À
ATOM	565	CB	ALA	173	-9.300	-1.725	1.292	1.00 12.08	A
ATOM	566	Ċ	ALA	173	-7.007	-2.537	0.653	1.00 15.86	A
		_	ALA	173	-6.147	-1.806	0.191		Ä
ATOM	567	0						1.00 14.20	
MOTA	568	N	GLN	174	-7.244	-3.714	0.109	1.00 16.56	A
MOTA	569	н	GLN	274	-7.774	-4.389	0.620	1.00 15.00	A
ATOM	570	CA	GLN	174	-6.470	-4.119	-1.070	1.00 19.25	A
ATOM	571	CB	GLN	174	-5.582	-5.292	-0.832	1.00 21.99	A
ATOM	572	ĊĠ	GLN	174	-4.205	-4.727	-1.030	1.00 30.99	A
	573	_	GLN	174	-3.174	-5.845			Ä
ATOM		CD					-0.979	1.00.34.25	
ATOM	574		GLN	174	-2.308	-5.899	-0.105	1.00 32.91	A
ATOM	575	NE2	GLN	174	-3.268	-6.699	-2.014	1.00 31.50	A
MCTA	576	HE21	GLN	174	-2.66B	-7.487	-1.970	1.00 15.00	A
MOTA	577	HE22	GLN	174	-3.973	-6.621	-2.714	1.00 15.00	A
ATOM	578	C	GLN	174	-7.413	-4.644	-2.114	1.00 19.20	À
ATOM	579	ō	GLN	-74	-8.285	-5.434	-1.880	1.00 20.03	A
ATOM	580	N	٧٨٠	175	.7.291	-4.107	-3.301	1.00 19.28	A
ATOM	581	H	VAL	175	- € . 594	-3.401	-3.400	1.00 15.00	A
MOTA	582	CA	VAL	175	-8.247	-4.500	-4.323	1.00 22.43	A
ATOM	583	CB	VAL	175	-9.319	-3.409	-4.644	1.00 21.41	- A
ATCM	584	CG1	VAL	175	-10.146	-2.830	-3.495	1.00 20.17	Ä
ATOM	585		VAL	175	-10.268	-4.061	-5.639	1.00 22.88	Ä
MCTA	586		VAL	175	-7.508		-, -	1.00 24.56	
		-							A
ATOM	587	0	VAL	:75	-6.928	-3.997	-6.301	1.00 23.28	A
ATOM	588	N	THR	176	-7.563	-6.180	-5.879	1.00 25.40	A
ATOM	589	н	THR	:76	-7.994	-6.850	-5.250	1.00 15.0G	A
ATOM	590	CA	THR	176	-7.086	-6.501	-7.222	1.00 24.46	A
ATOM	591	CB	THR	:76	-5.844	-7.454	-7.256	1.00 24.78	Ä
ATOM	592	061	THR	:76	-5.948	-8.650	-8.028	1.00 20.31	À
MOTA	593	H31	THR	. 176	-5.250	-9.253	-7.796	1.00 15.00	Ä
ATOM	594	232	THR	.76	·-£.329	-7.711	-5.867	1.00 17.37	÷
ATOM	595	Ξ	THR	176	-8.178	-6.700	-8.272	1.00 25.44	Ä
ATOM	596	Ĉ	THR	176	-9.326	-7.043	-7.995	1.00 26.86	Ä
ATOM	597		PHE	77	-7.855	-6.341	-9.506	1.00 22.44	Ä
				-77					•
ATOM	598		PHE		-6.920	-6.083	-9.732	1.00 15.00	÷
MOTA	599	CA	PHE	:77	-E.939	-6.511	-10.479	1.00 22.70	Ä

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FIGURE 1K

MCTA	600	5 5	PHE	177	- 5.746	-5.194	-10.599	1.00 20.90	÷
ATOM	501	CG	PHE	177	-8.813	-4.034	-10.927	1.00 11.E1	Ä
ATOM	602	<u> </u>	PHE	77	-8.771	-3.546	-12.252	1.00 00.11	Ä
MCTA	603	CD 2	PHE	277	-8.011	-3.422	-9.920	1.00 01.57	Ä
ATOM	504	CEI	PHE	77	-8.G41	-2.387	-12.550	1.00 20.53	÷
ATOM	605	CE2	PHE	177	-7.289	-2.247	-10.204	1.00 20.44	À
		cz	PHE	177	-7.376	-1.713	-11.500	1.00 22.79	À
ATOM	606		PHE	177	-8.381		-11.800	1.00 22.14	
ATOM	607	C	PHE	177	-7.219		-12.072	1.00 21.60	À
ATOM	608	0		178	-9.210		-12.625	1.00 24.52	Ä
ATOM	609	N	CYS	178	-10.146		-12.370	1.00 15.00	Ä
ATOM	610	H	CYS		-8.599			1.00 29.77	Ä
ATOM	611	CA	CYS	178	-8.501	-9.365	-14.214	1.00 32.06	Ä
ATOM	612	CB	CYS	178	-7.685		-15.792	1.00 35.17	Â
MOTA	613	SG	CYS	178	-9.323		-15.088	1.00 28.41	Ä
MOTA	614	C	CYS	178	-10.534		-15.185	1.00 27.54	À
ATOM	615	0	CYS	178	-8.589		-15.910	1.00 28.86	Ä
ATOM	616	N	SER	179			-15.754	1.00 15.00	Â
ATOM	617	H	SER	179	-7.608			1.00 13.00	À
ATOM	618	CA	SER	175	-9.374		-16.704		
MOTA	619	CB	SER	179	-9.379		-16.020	1.00 30.82	A
MOTA	620	OG	SER	179	-10.615		-16.319		A
MOTA	621	HG	SER	179	-10.725		-15.667	1.00 15.00	A
ATOM	622	-	SER	. 179	-9.063		-18.165	1.00 31.16	A
MOTA	623	0	SER	179	-7.931		-18.537	1.00 28.58	A
MOTA	624	N	ASN	180	-10.083		-19.042	1.00 35.32	A
MOTA	625	Н	ASN	180	-10.966		-18.834	1.00 15.00	A
MOTA	626	CA	ASN	180	-9.782		-20.366	1.00 34.74	A
ATOM	627	CB	ASN	180	-10.205		-21.589	1.00 37.96	A
ATOM	628	CG	ASN	180	-9.650		-22.896	1.00 37.12	A
ATOM	629	OD1	ASN	180	-10.058		-23.356	1.00 40.66	A
ATOM	630	ND2	ASN	180	-8.619		-23.456	1.00 35.85	A
MOTA	631	HD21	asn	180	-8.343		-23.306	1.00 15.00	A
ATOM	632	HD22	ASN	190	-8.153		-24.065	1.00 15.00	A
ATOM	633	C	ASN	180	-10.197		-20.588	1.00 36.96	A
MCTA	634	0	asn	180	-11.314	-2.894	-20.433	1.00 37.89	A
ATOM	635	N	arg	181	-9.147	-2.699		1.00 41.95	A
ATOM	636	H	ARG	181	-8.363	-3.318	-21.141	1.00 15.00	A
ATOM	637	CA	ARG	181	-8.997		-21.489	1.00 44.24	A
MOTA	638	CB	ARG	181	-7.563	-1.279	-22.026	1.00 43.43	A
MCTA	639	CG	arg	181	-6.34B	-1.638	-21.101	1.00 45.11	A
MOTA	640	CD	ARG	181	-6.235	-2.853	-20.134	1.00 40.68	A
ATOM	641	NE	ARG	181	-5.064	-2.772	-19.271	1.00 46.11	A
ATOM	642	HE	ARG	181	-4.991	-2.058	-18.578	1.00 15.00	A
ATOM	643	CZ	ARG	181	-4.024	-3.611	-19.432	1.00 49.77	À
MCTA	644	NH1	ARG	181	-2.886	-3.414	-18.790	1.00 54.33	Ä
MCTA	645	HH11	ARG	181	-2.113	-1.032	-18.918	1.00 15.00	À
ATOM	646	HH12		181	-2.807		-18.161	1.00 15.00	A
ATOM	647		ARG	181	-4.085		-20.247	1.00 54.26	A
ATOM	64B	HH21	ARG	181	-3.286		-20.354	1.00 15.00	A
MOTA	649		ARG	181	-4.918		-20.761	1.00 15.00	A
MOTA	650	Ξ	ARG	181	-10.049		-22.499	1.00 47.10	A
MCTA	651	၁	ARG	181	-10.979	-0.112	-22.227	1.00 49.20	Α.
ATOM	652	N	GLU	182	-9.895		-23.690	1.00 49.64	A
ATOM	653	H	GLU	182	-9.201	-2.166	-23.775	1.00 15.00	A
ATOM	€54	CA	32::	182	-10.976		-24.676	1.00 52.41	À
ATOM	655	CB	コニじ	182	-10.437		-25.970	1.00 56.93	Ä
MCTA	656	23	3೭೮	182	-10.932	-1.418	-27.295	1.00 66.05	A
MCTA	657	22	SLU	182	-10.758		-27.327	1.00 70.54	A
ATOM	658	CE:	SLU	162	-9.613			1.00 72.98	A
ATCM	65=		SLU	182	-11.778	0.830	-27.244	1.00 72.46	Ä

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FIGURE 1L

ATOM 661 D 3LU 182 -12.385 -1.534 -24.304 1:00 53.15 ATOM 662 N ALA 183 -12.505 -2.877 -23.215 1:00 54.15 ATOM 663 H ALA 183 -12.505 -2.877 -23.215 1:00 54.15 ATOM 664 CA ALA 183 -12.676 -3.173 -22.865 1:00 55.15 ATOM 665 CB ALA 183 -13.855 -3.258 -22.899 1:00 55.15 ATOM 666 C ALA 183 -13.855 -4.721 -22.447 1:00 45.02 ATOM 666 C ALA 183 -13.855 -4.721 -22.447 1:00 45.02 ATOM 667 O ALA 183 -15.712 -1.945 -21.990 1:00 47.77 ATOM 668 N SER 184 -13.773 -1.888 -20.878 1:00 52.95 ATOM 669 N SER 184 -12.826 -2.172 -20.991 1:00 15.00 ATOM 670 CA SER 184 -12.826 -2.172 -20.991 1:00 15.00 ATOM 671 CB SER 184 -13.384 -1.397 -18.481 1:00 53.58 ATOM 672 OG SER 184 -13.384 -1.397 -18.481 1:00 53.58 ATOM 673 NG SER 184 -13.291 -3.019 -17.388 1:00 15.00 ATOM 674 C SER 184 -13.291 -3.019 -17.388 1:00 15.00 ATOM 675 O SER 184 -13.291 -3.019 -17.388 1:00 15.00 ATOM 676 N SER 184 -13.291 -3.019 -17.388 1:00 15.00 ATOM 677 N SER 185 -14.623 0.345 -21.831 1:00 60.08 ATOM 678 CA SER 184 -13.291 -3.019 -17.388 1:00 15.00 ATOM 679 CB SER 184 -13.913 1:297 -18.964 1:00 65.25 ATOM 676 N SER 185 -14.623 0.345 -21.831 1:00 60.08 ATOM 677 N SER 185 -14.623 0.345 -21.831 1:00 60.08 ATOM 678 CA SER 185 -13.522 2.640 -22.869 1:00 60.49 ATOM 680 OG SER 185 -12.218 1:234 -22.833 1:00 15:00 ATOM 681 HG SER 185 -13.522 2.640 -22.869 1:00 60.49 ATOM 682 C SER 185 -14.623 0.345 -21.831 1:00 60.01 ATOM 683 C G GLN 186 -15.555 3.354 -17.050 8:00 59.95 ATOM 680 C G SER 185 -12.186 1:234 -22.833 1:00 15:00 ATOM 680 C G SER 185 -12.186 1:234 -22.833 1:00 15:00 ATOM 680 C G SER 185 -12.186 1:234 -22.833 1:00 15:00 ATOM 680 C G SER 185 -12.186 1:234 -22.833 1:00 15:00 ATOM 680 C G SER 185 -12.186 1:234 -22.833 1:00 15:00 ATOM 680 C G SER 185 -12.186 1:234 -22.833 1:00 15:00 ATOM 680 C G SER 185 -12.186 1:234 -22.833 1:00 15:00 ATOM 680 C G SER 185 -12.486 1:234 -22.833 1:00 15:00 ATOM 680 C G SER 185 -12.486 1:234 -22.833 1:00 15:00 ATOM 680 C G SER 186 -15.555 3:354 -17.050 1:00 59.59 ATOM 680 C G SER 186 -15.497 0:00 1:00 59.69 ATOM 680 C G SER 186 -15.4											
ATOM 662 N ALA 183 -11.676 -3.173 -22.862 1.00 51.31 ATOM 663 N ALA 183 -12.676 -3.173 -22.865 1.00 15.01 ATOM 664 CA ALA 183 -13.667 -3.1258 -22.899 1.00 15.01 ATOM 665 CB ALA 183 -13.667 -3.1258 -22.899 1.00 15.01 ATOM 665 CB ALA 183 -13.655 -4.721 -22.899 1.00 50.66 ATOM 666 C ALA 183 -14.662 -2.321 -21.867 1.00 50.66 ATOM 667 O ALA 183 -15.712 -1.845 -21.990 1.00 50.66 ATOM 668 N SER 184 -12.826 -2.172 -20.991 1.00 50.66 ATOM 669 H SER 184 -12.826 -2.172 -20.991 1.00 55.75 ATOM 670 CA SER 184 -12.826 -1.073 -7.19 1.00 55.78 ATOM 671 CB SER 184 -13.975 -2.448 1.7.721 1.00 47.46 ATOM 672 OG SER 184 -13.975 -2.448 1.7.721 1.00 47.46 ATOM 673 HG SER 184 -13.975 -2.448 1.7.721 1.00 47.46 ATOM 673 HG SER 184 -13.975 -2.448 1.7.721 1.00 47.46 ATOM 675 O SER 184 -13.971 1.07 17.9 880 1.00 59.95 ATOM 676 N SER 184 -13.973 -13.481 1.00 55.25 ATOM 676 N SER 184 -13.131 1.07 19.7 18.860 1.00 59.95 ATOM 676 N SER 185 -14.623 0.345 -21.331 1.00 60.08 ATOM 678 CA SER 185 -14.623 0.345 -21.331 1.00 60.12 ATOM 679 CB SER 185 -11.825 2.375 -21.391 1.00 60.12 ATOM 680 CG SER 185 -11.522 2.640 -22.869 1.00 60.49 ATOM 680 CG SER 185 -11.522 2.640 -22.869 1.00 60.49 ATOM 680 CG SER 185 -11.582 2.375 -21.331 1.00 15.00 ATOM 680 CG SER 185 -11.586 1.374 -22.833 1.00 15.00 ATOM 680 CG SER 185 -11.586 1.377 -19.670 1.00 59.95 ATOM 680 CG SER 185 -11.586 1.377 -19.670 1.00 59.95 ATOM 680 CG SER 185 -11.586 1.377 -19.670 1.00 59.95 ATOM 680 CG SER 185 -12.243 2.098 -23.242 1.00 59.80 ATOM 680 CG SER 185 -12.243 2.098 -23.242 1.00 59.80 ATOM 680 CG SER 185 -13.550 1.589 -20.885 1.00 59.59 ATOM 680 CG SER 185 -13.550 1.589 -20.885 1.00 59.59 ATOM 680 CG SER 185 -13.550 1.589 -20.885 1.00 59.59 ATOM 680 CG SER 185 -13.550 1.377 -1.00 59.60 ATOM 680 CG SER 185 -13.550 1.377 -1.00 59.60 ATOM 680 CG SER 185 -13.550 1.377 -1.00 59.60 ATOM 680 CG SER 185 -13.550 1.378 -1.00 59.95 ATOM 680 CG SER 185 -13.550 1.389 -20.885 1.00	OM.	660	2	3:::	152	-12.388	934	-24 304		23 00	
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ATOM 690 OE1 GLN 186 -17.270 3.513 -15.687 1.00 59.81 ATOM 691 NE2 GLN 186 -16.249 1.503 -15.787 1.00 59.63 ATOM 692 HE21 GLN 186 -16.249 1.503 -15.787 1.00 59.63 ATOM 693 HE22 GLN 186 -16.950 1.119 -15.168 1.00 15.00 ATOM 693 HE22 GLN 186 -16.950 1.119 -15.168 1.00 15.00 ATOM 694 C GLN 186 -14.758 6.290 -18.221 1.00 54.36 ATOM 695 O GLN 186 -15.596 7.198 -18.298 1.00 53.98 ATOM 696 N ALA 187 -13.566 6.424 -17.511 1.00 50.35 ATOM 697 H ALA 187 -13.566 6.424 -17.511 1.00 50.35 ATOM 698 CA ALA 187 -13.476 7.274 -16.970 1.00 15.00 ATOM 699 CB ALA 187 -12.388 5.599 -17.832 1.00 43.26 ATOM 699 CB ALA 187 -11.546 6.284 -18.918 1.00 38.95 ATOM 700 C ALA 187 -11.456 4.882 -16.849 1.00 40.48 ATOM 701 O ALA 187 -10.887 3.875 -17.295 1.00 43.24 ATOM 702 N PRO 188 -11.210 5.383 -15.594 1.00 38.66 ATOM 703 CD PRO 188 -11.210 5.383 -15.594 1.00 38.66 ATOM 704 CA PRO 188 -11.210 5.383 -15.594 1.00 38.66 ATOM 705 CB PRO 188 -10.840 3.783 -13.683 1.00 33.66 ATOM 706 CG PRO 188 -10.377 7.000 -14.036 1.00 33.99 ATOM 706 CG PRO 188 -10.840 3.783 -13.683 1.00 33.66 ATOM 707 C PRO 188 -10.840 3.783 -13.683 1.00 33.66 ATOM 708 O PRO 188 -10.840 3.783 -13.683 1.00 33.66 ATOM 709 N PHE 189 -10.147 2.695 -13.346 1.00 28.66 ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 28.66 ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 26.71 ATOM 712 CB PHE 189 -10.122 0.601 -12.034 1.00 26.71 ATOM 713 CG PHE 189 -10.122 0.601 -12.034 1.00 26.71 ATOM 714 CDI PHE 189 -10.122 0.601 -12.034 1.00 26.71 ATOM 715 CDI PHE 189 -10.122 0.601 -12.034 1.00 22.92 ATOM 716 CDI PHE 189 -10.122 0.601 -12.034 1.00 22.92 ATOM 716 CDI PHE 189 -10.122 0.601 -12.034 1.00 22.92 ATOM 716 CDI PHE 189 -10.122 0.601 -12.034 1.00 22.92 ATOM 716 CDI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 716 CDI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 716 CDI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 716 CDI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 716 CDI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 716 CDI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 716 CDI PHE 189 -10.59											´ A
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ATOM 692 HE21 GLN 186 -15.492 0.948 -16.113 1.00 15.00 ATOM 693 HE22 GLN 186 -16.950 1.119 -15.168 1.00 15.00 ATOM 695 C GLN 186 -14.758 6.290 -18.221 1.00 54.36 ATOM 695 O GLN 186 -15.596 7.198 -18.298 1.00 53.98 ATOM 696 N ALA 187 -13.566 6.424 -17.511 1.00 50.35 ATOM 697 H ALA 187 -13.566 6.424 -17.511 1.00 50.35 ATOM 698 CA ALA 187 -12.388 5.599 -17.832 1.00 43.26 ATOM 699 CB ALA 187 -12.388 5.599 -17.832 1.00 43.26 ATOM 699 CB ALA 187 -11.546 6.284 -18.918 1.00 38.95 ATOM 700 C ALA 187 -11.456 4.882 -16.849 1.00 40.48 ATOM 701 O ALA 187 -10.887 3.875 -17.295 1.00 43.24 ATOM 702 N PRO 188 -11.210 5.383 -15.594 1.00 38.66 ATOM 703 CD PRO 188 -11.210 5.383 -15.594 1.00 38.15 ATOM 704 CA PRO 188 -11.210 5.383 -15.594 1.00 38.15 ATOM 705 CB PRO 188 -10.220 4.665 -14.751 1.00 35.94 ATOM 706 CG PRO 188 -10.377 7.000 -14.036 1.00 33.99 ATOM 706 CG PRO 188 -10.377 7.000 -14.036 1.00 33.41 ATOM 708 O PRO 188 -10.840 3.783 -13.683 1.00 33.666 ATOM 708 O PRO 188 -10.840 3.783 -13.466 1.00 28.666 ATOM 708 O PRO 188 -10.840 3.783 -13.466 1.00 28.666 ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 15.00 ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 15.00 ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 26.71 ATOM 712 CB PHE 189 -10.122 0.601 -12.034 1.00 26.71 ATOM 713 CG PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.0849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.0849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.0849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.0849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.0849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.0849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.0849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.0811 1.00 21.13 ATOM 7					186	-17.270	3.513	-15.687	1.00	59.81	A
ATOM 693 HE22 GLN 186 -16.950 1.119 -15.168 1.00 15.00 ATOM 694 C GLN 186 -14.758 6.290 -18.221 1.00 54.36 ATOM 695 O GLN 186 -15.596 7.198 -18.298 1.00 53.98 ATOM 696 N ALA 187 -13.566 6.424 -17.511 1.00 50.35 ATOM 697 H ALA 187 -13.476 7.274 -16.970 1.00 15.00 ATOM 698 CA ALA 187 -12.388 5.599 -17.832 1.00 43.26 ATOM 699 CB ALA 187 -11.546 6.284 -18.918 1.00 38.95 ATOM 700 C ALA 187 -11.456 4.882 -16.849 1.00 40.48 ATOM 701 O ALA 187 -10.887 3.875 -17.295 1.00 40.48 ATOM 702 N PRO 188 -11.210 5.383 -15.594 1.00 38.66 ATOM 703 CD PRO 188 -11.543 6.687 -15.000 1.00 38.15 ATOM 704 CA PRO 188 -11.543 6.687 -15.000 1.00 38.15 ATOM 705 CB PRO 188 -9.395 5.813 -14.150 1.06 33.99 ATOM 706 CG PRO 188 -10.377 7.000 -14.036 1.00 33.41 ATOM 707 C PRO 188 -10.377 7.000 -14.036 1.00 33.41 ATOM 708 O PRO 188 -10.840 3.783 -13.683 1.00 33.66 ATOM 708 O PRO 188 -10.840 3.783 -13.683 1.00 33.41 ATOM 709 N PHE 189 -10.147 2.695 -13.346 1.00 28.66 ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 15.00 ATOM 711 CA PHE 189 -10.122 0.601 -12.034 1.00 26.71 ATOM 712 CB PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 713 CG PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD PHE 189 -10.571 -0.852 1.00 19.12 ATOM 715 CD PHE 189 -10.571 -1.806 -8.736 1.00 18.44	DM 6	691	NE2	GLN	186	-16.249	1.503	-15.787	1.00	59.63	A
ATOM 694 C GLN 186 -14.758 6.290 -18.221 1.00 54.36 ATOM 695 O GLN 186 -15.596 7.198 -18.298 1.00 53.98 ATOM 696 N ALA 187 -13.566 6.424 -17.511 1.00 50.35 ATOM 697 H ALA 187 -13.476 7.274 -16.970 1.00 15.00 ATOM 698 CA ALA 187 -12.388 5.599 -17.832 1.00 43.26 ATOM 699 CB ALA 187 -11.546 6.284 -18.918 1.00 38.95 ATOM 700 C ALA 187 -11.456 4.882 -16.849 1.00 40.48 ATOM 701 O ALA 187 -10.887 3.875 -17.295 1.00 43.24 ATOM 702 N PRO 188 -11.210 5.383 -15.594 1.00 38.66 ATOM 703 CD PRO 188 -11.543 6.687 -15.000 1.00 38.15 ATOM 704 CA PRO 188 -11.543 6.687 -15.000 1.00 38.15 ATOM 705 CB PRO 188 -9.395 5.813 -14.150 1.06 33.99 ATOM 706 CG PRO 188 -9.395 5.813 -14.150 1.06 33.99 ATOM 707 C PRO 188 -10.377 7.000 -14.036 1.00 32.69 ATOM 708 O PRO 188 -10.840 3.783 -13.683 1.00 33.666 ATOM 709 N PHE 189 -10.147 2.695 -13.346 1.00 28.66 ATOM 709 N PHE 189 -10.147 2.695 -13.346 1.00 28.66 ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 15.00 ATOM 711 CA PHE 189 -10.122 0.601 -12.034 1.00 26.71 ATOM 712 CB PHE 189 -10.671 -0.189 10.849 1.00 22.92 ATOM 713 CG PHE 189 -10.671 -0.189 10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.571 -0.895 -10.011 1.00 21.13 ATOM 715 CDI PHE 189 -10.571 -1.806 -8.736 1.00 18.44)M 6	692	HE21	GLN	186	-15.492	0.948	-16.113	1.00	15.00	A
ATOM 694 C GLN 186 -14.758 6.290 -18.221 1.00 54.36 ATOM 695 O GLN 186 -15.596 7.198 -18.298 1.00 53.98 ATOM 696 N ALA 187 -13.566 6.424 -17.511 1.00 50.35 ATOM 697 H ALA 187 -13.476 7.274 -16.970 1.00 15.00 ATOM 698 CA ALA 187 -12.388 5.599 -17.832 1.00 43.26 ATOM 699 CB ALA 187 -11.546 6.284 -18.918 1.00 38.95 ATOM 700 C ALA 187 -11.456 4.882 -16.849 1.00 40.48 ATOM 701 O ALA 187 -10.887 3.875 -17.295 1.00 43.24 ATOM 702 N PRO 188 -11.210 5.383 -15.594 1.00 38.66 ATOM 703 CD PRO 188 -11.543 6.687 -15.000 1.00 38.15 ATOM 704 CA PRO 188 -10.220 4.665 -14.751 1.00 35.94 ATOM 705 CB PRO 188 -9.395 5.813 -14.150 1.06 33.99 ATOM 706 CG PRO 188 -10.377 7.000 -14.036 1.00 33.66 ATOM 708 O PRO 188 -10.377 7.000 -14.036 1.00 33.41 ATOM 709 N PHE 189 -10.147 2.695 -13.346 1.00 33.41 ATOM 709 N PHE 189 -9.260 2.508 -13.748 1.00 15.00 ATOM 711 CA PHE 189 -9.260 2.508 -13.748 1.00 26.21 ATOM 712 CB PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 713 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 714 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CDI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 715 CDI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 715 CDI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 715 CDI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 715 CDI PHE 189 -11.571 -1.806 -8.736 1.00 18.44	M 6	593	HE22	GLN	186	-16.950	1.119	-15.168	1.00	15.00	A
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ATOM 708 O PRO 188 -11.885 4.062 -13.140 1.00 33.41 ATOM 709 N PHE 189 -10.147 2.695 -13.346 1.00 28.66 ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 15.00 ATOM 711 CA PHE 189 -10.721 2.013 -12.171 1.00 26.71 ATOM 712 CB PHE 189 -10.122 0.601 -12.034 1.00 26.21 ATOM 713 CG PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 714 CD1 PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 715 CD2 PHE 189 -10.126 0.005 -9.566 1.00 17.72 ATOM 715 CD2 PHE 189 -11.687 -1.165 -11.064 1.00 21.88 ATOM 715 CE1 PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 715 CE2 PHE 189 -12.124 -1.995 -10.011 1.00 21.13 ATOM 715 CE2 PHE 189 -11.571 -1.806 -8.736 1.00 18.44				PRO					1.00	32.69	A
ATOM 709 N PHE 189 -10.147 2.695 -13.346 1.00 28.66 ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 15.00 ATOM 711 CA PHE 189 -10.721 2.013 -12.171 1.00 26.71 ATOM 712 CB PHE 189 -10.122 0.601 -12.034 1.00 26.21 ATOM 713 CG PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 714 CD1 PHE 189 -10.126 0.005 -9.566 1.00 17.72 ATOM 715 CD2 PHE 189 -11.687 -1.165 -11.064 1.00 21.88 ATOM 716 CE1 PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 717 CE2 PHE 189 -12.124 -1.995 -10.011 1.00 21.13 ATOM 718 CC PHE 189 -11.571 -1.806 -8.736 1.00 18.44			С			-10.840	3.783	-13.683	1.00	33.66	А
ATOM 709 N PHE 189 -10.147 2.695 -13.346 1.00 28.66 ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 15.00 ATOM 711 CA PHE 189 -10.721 2.013 -12.171 1.00 26.71 ATOM 712 CB PHE 189 -10.122 0.601 -12.034 1.00 26.21 ATOM 713 CD PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 714 CDI PHE 189 -10.126 0.005 -9.566 1.00 17.72 ATOM 715 CDI PHE 189 -11.687 -1.165 -11.064 1.00 21.88 ATOM 716 CEI PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 717 CEI PHE 189 -12.124 -1.995 -10.011 1.00 21.13 ATOM 718 CD PHE 189 -11.571 -1.806 -8.736 1.00 18.44)M 7	708	0	PRO	188	-11.885	4.062	-13.140	1.00	33.41	A
ATOM 710 H PHE 189 -9.260 2.508 -13.748 1.00 15.00 ATOM 711 CA PHE 189 -10.721 2.013 -12.171 1.00 26.71 ATOM 712 CB PHE 189 -10.122 0.601 -12.034 1.00 26.21 ATOM 713 CG PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 714 CD1 PHE 189 -10.126 0.005 -9.566 1.00 17.72 ATOM 715 CD2 PHE 189 -11.687 -1.165 -11.064 1.00 21.88 ATOM 716 CE1 PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 717 CE2 PHE 189 -12.124 -1.995 -10.011 1.00 21.13 ATOM 718 CD PHE 189 -11.571 -1.806 -8.736 1.00 18.44	M 7	705	N	PHE	189	-10.147	2.695	-13.346			, A
ATOM 711 CA PHE 189 -10.721 2.013 -12.171 1.00 26.71 ATOM 712 CB PHE 189 -10.122 0.601 -12.034 1.00 26.21 ATOM 713 CG PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 714 CD1 PHE 189 -10.126 0.005 -9.566 1.00 17.72 ATOM 715 CD2 PHE 189 -11.687 -1.165 -11.064 1.00 21.88 ATOM 716 CE1 PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 717 CE2 PHE 189 -12.124 -1.995 -10.011 1.00 21.13 ATOM 718 CC PHE 189 -11.571 -1.806 -8.736 1.00 18.44	OM 7	710	н	PHE							À
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ATOM 713 CG PHE 189 -10.671 -0.189 -10.849 1.00 22.92 ATOM 714 CD1 PHE 189 -10.126 0.005 -9.566 1.00 17.72 ATOM 715 CD2 PHE 189 -11.687 -1.165 -11.064 1.00 21.88 ATOM 716 CE1 PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 717 CE2 PHE 189 -12.124 -1.995 -10.011 1.00 21.13 ATOM 718 CZ PHE 189 -11.571 -1.806 -8.736 1.00 18.44											
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ATOM 715 CD2 PHE 189 -11.687 -1.165 -11.064 1.00 21.88 ATOM 716 CE1 PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 717 CE2 PHE 189 -12.124 -1.995 -10.011 1.00 21.13 ATOM 718 CZ PHE 189 -11.571 -1.806 -8.736 1.00 18.44		- • -			. 0 5						A
ATOM 716 CE1 PHE 189 -10.590 -0.815 -8.522 1.00 19.12 ATOM 717 CE2 PHE 169 -12.124 -1.995 -10.011 1.00 21.13 ATOM 718 CE PHE 189 -11.571 -1.806 -8.736 1.00 18.44)M =		~~-								Č
ATOM 717 CEL PHE 169 -12.124 -1.995 -10.011 1.00 21.13 ATOM 718 CE PHE 189 -11.571 -1.806 -8.736 1.00 18.44	M 1										A
ATOM 718 CC PHE 189 -11.571 -1.806 -8.736 1.00 18.44	JY:	5									A
ALOM 115 LD PHE 189 -11.571 -1.806 -8.736 1.00 18.44											A
			===								A
ATOM	ייינע. דיינע		-	PHE	189	-10.445	2.815	-10.909	1.00	27.14	A

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FIGURE 1M

ATOM	72:	5	PHE	189	-9.308	3.244	-10.706	: . : : : : : : : : : : : : : : : : : :	À
	72:	:	ΞLΕ	195	-11.469	2.964	-::::	1.00 04 71	À
ATOM	722	Ä	===	190	-12.408		-10.388	1.11 15.11	÷.
ATOM		ĊA	ΞΞĒ	190	-11.193	3.626	-8.788	1.00 14.03	À
ATOM	723		ΞĒ	193	-11.316	5.242	-8.743	1.00 16.86	Ä
MCTA	724	CE .			-11.892	5.979	-9.997	1.00 19.87	Ä
MOTA	725	CS2	ILE	190	-11.801			1.00 23.54	Ä
MOTA	726	CGI	ILE	190		5.888	-7.424		Ä
ATOM.	727	557	ILE	190	-12.819	7.012	-7.645		
ATOM	728	\subseteq	ILE	190	-11.844	2.812	-7.656	1.00 21 97	÷
ATOM	729	0	ILE	190	-12.891	2.197	-7.801	1.00 16.30	÷
ATOM	73C	N	ALA	191	-11.026	2.700	-6.590	1.00 17.21	A
ATOM	731	H	ALA	191	-10.124	3.124	-6.662	1.00 15.00	Ä
ATOM	732	CA	ALA	191	-11.501	2.195	-5.321	1.00 15.20	۸
ATOM	733	CB	ALA	191	-10.730	0.928	-4.968	1.00 14.79	Ä
	734	c	ALA	191	-11.439	3.230	-4.206	1.00 17.11	À
ATOM	735	Ö	ALA	191	-10.467	3.961	-4.052	1.00 14.04	۸
ATOM		N	SER	192	-12.511	3.245	-3.433	1.00 14.72	À
ATOM	736			192	-13.277	2.694	-3.804	1.00 15.00	A
ATOM	737	H	SER		-12.725	4.289	-2.423	1.00 16.69	A
ATOM	738	CA	SER	192		5.144	-2.803	1.00 14.83	Ā
MOTA	739	CB	SER	192	-13.931			1.00 21.23	Ä
MOTA	740	OG	SER	192	-13.556	5.828	-3.994	•	Â
ATOM	741	HG	SER	192	-14.367	5.966	-4.520	1.00 15.00	
ATOM	742	С	SER	192	-12.980	3.682	-1.069	1.00 17.77	Ä
ATOM	743	0	SER	192	-13.753	2.738	-0.947	1.00 20.76	A
ATOM	744	N	LEU	193	-12.285	4.209	-0.038	1.00 15.56	A
ATOM	745	H	LEU	193	-11.681	4.959	-0.280	1.00 15.00	À
ATOM	746	CA	LEU	193	-12.510	3.761	1.366	1.00 13.27	A
ATOM	747	CB	LEU	193	-11.195	3.825	2.217	1.00 12.74	À
ATOM	748	ČŠ	LEU	193	-11.051	3.141	3.604	1.00 14.37	À
	749	CD1	LEU	193	-12.272	2.354	4.116	1.00 14.67	A
ATOM	750	CD2	LEU	193	-10.274	3.986	4.622	1.00 12.64	A
ATOM				193	-13.497	4.748	1.911	1.00 11.22	A
MCTA	751	2	LEU		-13.188	5.912	1.903	1.00 12.22	Ä
ATOM	752	C	LEU	193		4.326	2.310	1.00 13.66	A
ATOM	753	N	CYS	194	-14.652	3.347	2.276	1.00 15.00	Ä
MCTA	754	H	CYS	194	-14.828		2.713	1.00 14.84	Ä
ATOM	755	CA	CYS	194	-15.595	5.360		1.00 17.58	Ä
ATOM	756	CB	CY5	194	-16.915	5.409	1.918		Ä
ATOM	757	SG	CYS	194	.16.623	5.417	0.165	1.00 16.33	
ATOM	758	C	CYS	194	-16.046	5.163	4.137	1.00 12.81	A
ATOM	759	0	CYS	194	-15.983	4.072	4.655	1.00 10.34	A
ATOM	760	N	LEU	195	-16.557	6.254	4.697	1.00 14.32	Ä
ATOM	761	H	LEU	195	-16.541	7.088	4.154	1.00 15.00	A
ATOM	762	CA	LEU	:95	-17.039	6.291	6.076	1.00 14.89	, A
ATOM	763	CB	LEU	195	-16.195	7.372	6.789	1.00 15.56	A
ATOM	764	CG	LEU	195	-16.571	7.680	8.242	1.00 15.56	À
MCTA	765	55:	LEU	:95	-15.932	8.967	8.762	1.00 13.72	À
ATOM	765	CD2		:95	-16.463	6.448	9.154	1.00 17.25	A
ATOM	757	5	LEU	195	-19.546	6.544	6.209	1.00 13.54	A
ATOM	768	ò	LEU	195	-19.038	7.521	5.705	1.00 14.56	A
			LYS	196	.19.238	5.667	6.905	1.00 16.36	Ä
ATOM	769	N		196	-18.719	4.875	7.197	1.00 15.00	A
ATOM	770	H	LYS		-20.577	5.972	7.405	1.00 21.01	Ä
MOTA	771	CA	LYS	196				1.00 22.66	Ä
ATOM	772	CB	LYS	196	-21.475	4.726	7.146		· •
ATOM	773	23	LYS	196	-22.953	4.839	7.590	1.00 31.25	Ä
ATOM	;	==	LYS	: • •	-23.364	4.915	9.104	1.00 40.25	•
ATOM	5	ΞΞ	LYS	. 9 ÷	-23.189	3.694	10.060	1.00 43.56	÷
ATOM	776	:::	:.YS	196	-23.604	4.158	11.453	1.00 44.46	Ţ
ATOM		HZ:	LYS	196	-22.182	4.799	11.467	1.00 15.00	Ä
ATOM	~~ <u> </u>	HZ	LYS	19€	-23.847	4.665	11.778	1.00 15.00	Ä
ATOM	~~ =		LYS	196	-22.857	3.334	12.966	1.00 15.00	Ä
	•								

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FIGURE 1N

ATOM	780	5	LYS	196	-20.478	6.290	8.899	1.00 19.25	Ä
ATOM	781	C	LYS	196	-20.194	5.434	9.714	1.00 18.38	Ä
ATOM	782	N	SER	197	-20.664	7.534	9.272	1.00 20.63	.
ATOM	753	Ħ	SER	197	-20.891	5.247	6.615	1.22 18.15	÷.
ATOM	784	CA	SER	197	-20.752	7.701	10.729	1.00 24.87	Ä
ATOM	795	CB	SER	197	-19.898	8.878	11.207	1.00 24.87	Ê
ATOM	786	OG	SER	267	-19.563	9.687	12.568	1.00 32.22	
	787	23	SER	197	-18.795	8.110			À
ATOM							12.611	1.00 15.00	A
ATOM	788	C	SER	197	-22.216	7.810	11.219	1.00 26.33	À
ATOM	789	0	SER	197	-23.078	8.303	10.497	1.00 26.57	Ä
ATOM	790	N	PRO	198	-22.534	7.274	12.407	1.00 25.77	Ä
ATOM	791	CD	PRO	198	-21.649	6.526	13.351	1.00 32.92	À
ATOM	792	CA	PRC	198	-23.919	7.381	12.913	1.00 26.73	Ä
ATOM	793	CB	PRO	198	-23.784	6.789	14.318	1.00 32.89	A
ATOM	794	CG	PRO	198	-22.289	6.726	14.659	1.00 33.55	Ä
ATOM	795	C	PRO	198	-24.591	8.789	12.847	1.00 26.60	À
ATOM	796	0	PRO	198	-24.035	9.817	13.242	1.00 20.20	Ä
ATOM	797	N	GLY	199	-25.729	8.773	12.119	1.00 25.75	Ä
ATOM	798	Н	GLY	199	-26.170	7.857	12.057	1.00 15.00	Ä
ATOM	799	CA	GLY	199	-26.486	10.003	11.790	1.00 26.91	Â
ATOM	800	Ć	GLY	199	-25.821	10.003			
							10.816	1.00 28.98	A
ATOM	801	0	GLY	199	-26.084	12.151	10.797	1.00 31.05	Α,
ATOM	802	N	ARG	200	-24.898	10.464	10.001	1.00 30.15	λ
ATOM	803	H	ARG	200	-24.629	9.519	10.165	1.00 15.00	À
MOTA	804	CA	ARG	200	-24.140	11.384	9.166	1.00 28.98	A
ATOM	805	CB	arg	200	-22.749	11.590	9.783	1.00 33.16	A
ATOM	806	CG	ARG	200	-22.739	12.290	11.162	1.00 38.34	A
ATOM	807	CD	ARG	200	-21.327	12.530	11.705	1.00 42.14	A
ATOM	808	NE	ARG	200	-21.292	12.875	13.131	1.00 43.64	Ä
ATOM	809	HE	ARG	200	-21.327	13.831	13.424	1.00 15.00	A
ATOM	810	CZ	ARG	200	-21.138	11.896	14.051	1.00 46.40	Ä
ATOM	811	NH1	ARG	200	-21.219	10.603	13.733	1.00 46.31	Ä
ATCM	812	HH11	ARG	206	-21.104	9.910	14.445	1.00 15.00	Ä
ATOM	813	HH12		200	-21.394	10.320	12.789	1.00 15.00	Ä
ATOM	814		ARG	200	-20.901	12.226	15.311	1.00 46.65	Ä
ATOM		HH21	ARG	200	-20.847	13.193	15.566	1.00 15.00	
ATOM	816	HH22		200	-20.785				Ą
MOTA	817		ARG			11.510	16.002	1.00 15.00	À
		2		20C	-24.084	10.967	7.710	1.00 27.77	À
ATOM	818	C	ARG	200	-24.264	9.791	7.449	1.00 28.21	Ä
ATOM	819	N	PHE	201	-23.853	11.926	6.792	1.00 30.83	Ä
MOTA	820	H	PHE	201	-23.513	12.821	7.126	1.00 15.00	A
ATOM	821	CA	PHE	201	-24.016	11.70B	5.339	1.00 34.17	A
MCTA	922	CB	PHE	201	-23.851	12.996	4.572	1.00 31.58	A
ATOM	823	CG	PHE	201	-25.154	13.730	4.614	1.00 34.85	Α
MCTA	524	CD1	PHE	201	-25.174	15.062	5.081	1.00 37.56	A
MCTA	825	CD2	PHE	201	-26.335	13.081	4.190	1.00 37.89	Ä
MOTA	82€	CEl	PHE	201	-26.397	15.749	5.182	1.00 36.91	A
MOTA	827	CE2	PHE	201	-27.566	13.762	4.280	1.00 38.98	Ä
MOTA	828	CZ	PHE	201	-27.572	15.065	4.815	1.00 37.61	A
ATOM	829	Ξ	PHE	201	-23.277	10.605	4.545	1.00 39.40	Ä
MCTA	83C	C	PHE	201	-23.853	10.034	3.604	1.00 45.71	Ä
ATOM	831	N	GLU	202	-22.031	10.316	5.034	1.00 35.75	
MCTA	832	H	GLU	202	-21.878	10.753	5.925		Ä
ATOM	833	 Ea	320	202	-20.964	9.564	4.318		÷
ATOM	834	CE	32:	212	-21.295			1.00 34.52	÷
ATOM	835	-= ::3	27	252		8.540	3.234	1.00 33.66	÷
ATOM			 -		-21.924	7.245	3.713	1.00 40.61	Ä
	83 £	22	320	202	-22.647	6.505	2.561	1.00 46.12	Ä
ATOM	837	CE:	27	202	-23.461	5.613	2.886	1.00 45.89	A
ATOM	836	CES	310	202	-22.417	6.814	1.370	1.00 45.63	Ä
ATCM	639	0	SLU	252	-19.924	10.450	3.717	1.00 29.99	÷

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FIGURE 10

ATOM	940 0 3	ilu ici	-20.137	11.567	3.300	1.00 30.76	A
	•	JRS 203	-15.728	9.897	3.556	1.00 26.88	÷.
ATOM			-18.690	8.998	4.285	1.00 15.00	Ä
ATOM	~		-17.539	10.603	3.355	1.00 18.00	À
MCTA		RG 203	-16.819	11.410	4.457	I.00 IT.07	Ä
MCTA	• • •	JRG 213				1.00 37.32	Ä
MCTA	845 CG A	\RG 203	-17.681	12.187	5.467		Â
ATOM	846 CD A	LRG 203	-16.894	13.213	6.339	1.00 48.09	Ţ
ATOM		JRG 203	-15.911	12.667	7.308	1.00 56.90	Ä
	• • • • • •	IRG 203	-16.240	12.433	8.223	1.00 15.00	À.
ATOM		IRG 203	-14.572	12.475	7.001	1.00 66.77	À
MOTA		RG 203	-13.702	12.002	7.911	1.00 68.44	À
MOTA			-12.745	11.829	7.666	1.00 15.00	À
MOTA			-14.016	11.822	8.845	1.00 15.00	A
ATOM		ARG 203			5.766	1.00 67.68	À
ATOM	853 NH2 A	ARG 203	-14.084	12.716		1.00 15.00	Ä
ATOM	854 HH21 A	URG 203	-14.670	13.108	5.060		Ä
ATOM	855 HH22 A	LRG 203	-13.143	12.499	5.544	1.00 15.00	
ATOM	856 C A	ARG 203	-16.517	9.633	2.678	1.00 17.71	À
ATOM		ARG 203	-16.375	9.418	2.931	1.00 7.69	À
	• -	LE 204	-15.789	10.253	1.791	1.00 14.42	À
ATOM		LE 204	-15.915	11.228	1.561	1.00 15.00	A
ATOM			-14.662	9.482	1.353	1.00 18.32	À
ATOM		LE 204	-14.520	9.392	-0.231	1.00 24.52	Ä
ATOM		LE 204		9.529	-1.069	1.00 21.85	A
MOTA		ILE 204	-15.820		-0.949	1.00 26.35	A
ATOM		ILE 204	-13.439	10.195		1.00 36.33	A
ATOM		ILE 204	-13.992	11.231	-1.961	1.00 16.58	Ä
MOTA		ILE 204	-13.387	9.819	2.153		Ä
ATOM	866 C 1	ILE 204	-13.070	10.956	2.457	1.00 18.63	
MOTA	867 N I	LEU 205	-12.718	8.725	2.571	1.00 13.32	Ä
ATOM	868 H I	LEU 205	-13.142	7.853	2.321	1.00 15.00	A
ATOM		LEU 205	-11.467	8.829	3.322	1.CO 10.01	A
MCTA		LEU 205	-11.440	7.688	4.382	1.00 6.66	A
		LEU 205	-12.571	7.727	5.441	1.00 7.99	A
MCTA			-12.722	9.088	6.089	1.00 8.78	A
MOTA			-12.419	6.720	6.582	1.00 8.08	A
ATOM		LEU 205	-10.268	8.811	2.377	1.00 9.75	A
ATCM	• .		-9.416	9.655	2.320	1.00 10.25	A
ATOM	. •		-10.252	7.769	1.562	1.00 10.28	A
ATOM		LEU 206		7.119	1.684	1.00 15.00	Α
ATOM	· ·	LEU 206	-10.991	_	0.610	1.00 10.02	A
ATOM		LEU 206	-9.166	7.555		1.00 11.94	A
ATOM		LEU 206	-8.249	6.384	0.990	-	Ä
ATOM		LEU 206	-7.001	6.527	1.859	1.00 14.40	Â
ATOM	881 CD1	LEU 206	-7.094	5.595	3.074	1.00 14.49	Â
ATOM	882 CD2	LEU 206	-€.531	7.958	2.151	1.00 8.78	
MCTA	883 C	LEU 206	· 9 . 756	7.071	-0.697	1.00 11.91	A
ATOM	884 C	LEU 206	-10.792	6.406	-0.778	1.00 10.67	Ą
ATOM		ARG 207	-9.005	7.428	-1.720	1.00 8.06	A
ATOM		ARG 207	-8.196	7.992	-1.553	1.00 15.00	Ä
ATOM		ARG 207	-9.309	6.823	-2.992	1.00 10.45	A
ATOM		ARG 207	-9.974	7.790	-3.904	1.00 8.71	Ä
		ARG 207	-11.258	8.270	-3.357	1.00 15.68	A
ATOM	_	ARG 207	-11.652	9.459	-4.163	1.00 22.25	A
ATOM			-12.670	9.192	-5.171	1.00 29.59	A
ATOM			-13.115	8.300	-5.249	1.00 15.00	A
ATOM	892 HE	ARG 207				1.00 40.09	Ä
MCTA	993 CZ	ARG 207	-13.063	10.272	-5.919	1.00 40.09	Ä
ATOM	894 NH1	AKJ	-12.482	11.498	-5.813		Ä
ATOM	895 HHII	ARS 207	-12.813	12.246	-6.391	1.00 15.00	, ,
ATOM	ife HHLD	ARG 207	-11.737	11.651	-5.165	1.00 15.00	
MOTA	897 NH2	ARG 207		10.111	-6.773	1.00 40.86	
ATOM	ese HHZI	ARG 207	:4 392	10.877	-7.329	1.00 15.00	Ä
ATOM	SSS HHIL		-14.498	9.207	-6.853	1.00 15.00	Ä

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FIGURE 1P

ATOM	900	<u>-</u>	ARG	257	-B.044	6.456	-3.741	1.00 10.59	À
ATOM	901	Š	ARG	207	-7.053	7.153	-3.787	1.00 15.55	÷.
ATOM	902	N	شنة	208	-8.396	5.358	-4.465	1.00 1 ⁴ .04	A
ATOM	903	H	٨٠٠٨	208	-8.875	4.758	-4.355	1.00 15.00	A
	904	CA	٨٠٠٨	209	-7.925	5.128	-5.465	1.00 17.00	À
ATOM	905	CB	ALA	206	-6.052	4.020	-5.072	1.00 14.69	À
MCTA		C	ALA	208	-7.544	4.830	-6.854	1.00 20.46	· A
ATOM	906	0	ALA	208	-8.438	4.020	-7.057	1.00 21.89	Ä
ATOM	907		مسم هنته	203	-6.986	5.586	-7.808	1.00 26.22	Ä
ATOM	908	N H	ALA	209	-6.280	6.235	-7.533	1.00 15.00	Ä
ATOM	909			209	-7.253	5.208	-9.196	1.00 28.06	Ä
ATOM	910	CA	ALA		-7.702	6.380	-10.069	1.00 27.10	Ä
ATOM	911	CB	ALA	209			-9.832	1.00 32.54	2
ATOM	912	C	ALA	209	-6.075	4.461		1.00 33.00	Ä
ATOM	913	0	ALA	209	-4.895	4.726	-9.593		Ä
MOTA	914	N	ASN	210	-6.502	3.491	-10.634		
MOTA	915	H	ASN	210	-7.466	3.249	-10.531	1.00 15.00	A
ATOM	916	CA	ASN	210	-5.674	2.893	-11.662	1.00 36.00	À
ATOM	917	CB	ASN	210	-5.366	1.446	-11.355	1.00 39.53	A
ATOM	918	CG	ASN	210	-4.463		-10.154	1.00 42.59	Ä
MOTA	919	OD1	ASN	210	-4.285	2.273	-9.342	1.00 39.26	À
MOTA	920	ND2		210	-3.951	-	-10.055	1.00 41.77	A
ATOM	921	HD21	ASN	210	-3.990	-0.479	-10.817	1.00 15.00	A
MCTA	922	HD22	ASN	210	-3.364	-0. 08 1	-9.279	1.00 15.00	A
ATOM	923	С	ASN	210	-6.299	2.931	-13.043	1.00 36.95	A
ATOM	924	0	ASN	210	-7.492	2.752	-13.259	1.00 36.93	A
ATOM	925	N	THR	211	-5.447	3.168	-14.013	1.00 37.83	A
ATOM	926	н	THR	211	-4.484	3.377	-13.821	1.00 15.00	A
ATOM	927	CA	THR	211	-6.119	3.224	-15.314	1.00 41.27	A
ATOM	928	CB	THR	211	-5.325	4.158	-16.268	1.00 44.53	A
ATOM	929	OG1	THR	211	-6.076	4.506	-17.438	1.00 49.34	A
MCTA	930	HG1	THR	211	-€.032	5.493	-17.508	1.00 15.00	A
MCTA	931	CG2	THR	211	-3.926	3.604	-16.581	1.00 46.08	A
ATOM	932	c	THR	211	-6.434	1.833	-15.87B	1.00 39.17	A
ATOM	933	ō	THR	211	-5.822	0.863	-15.475	1.00 36.48	A
ATOM	934	N	HIS	212	-7.416	1.718	-16.789	1.00 37.14	Ä
	935	н	HIS	212	-8.106	2.438	-16.87B	1.00 15.00	A
ATOM	936	CA	HIS	212	-7.294	0.454	-17.529	1.00 33.23	A
MOTA			HIS	212	-8.680	-0.012	-18.082	1.00 27.73	A
ATOM	937	CB CG	HIS	212	-9.856	0.060	-17.111	1.00 24.58	A
ATOM	938		HIS	212	-10.862	0.967	-17.161	1.00 24.59	A
ATOM	939		HIS	212	-11.000	1.702	-17.794	1.00 15.00	A
ATOM	940				-10.049	-0.723	-15.985	1.00 20.65	Ä
MOTA	941		HIS	212		-0.723	-15.383	1.00 24.01	Ä
ATOM	942	NE2		212	-11.154	0.780	-16.092	1.00 17.59	Ä
ATOM	943	CEI	HIS	2:2	-11.665		-18.683	1.00 38.31	Ä
ATOM	944	Č	HIS	212	-6.257	0.633		1.00 33.92	Ä
ATOM	945	0	415	212	-5.363	-6.132	-18.923		Ä
ATOM	946	N	SER	213	-6.444		-19.443	1.00 46.63	Â
ATOM	947	H	SER	213	-7.156		-19.055		Â
atom	948	CA	SER	213	-5.705		-20.675	1.00 53.91	Â
ATOM	949	CB	SER	213	-4.272	2.704	-20.400	1.00 52.61	Â
ATOM	950	OG	SER	213	-3.266		-20.547	1.00 53.97	•
MOTA	951	HG	SER	213	-3.363	1.064	-19.823	1.00 15.00	Ä
ATOM	952	C	SER	213 213	-5.844		-22.097	1.00 60.03	A
ATOM	953	0	SER	213	-5.005		-22.682	1.00 61.19	ÿ
ATOM	954	×	SER	214	-7.043	1.803		1.00 64.96	À
ATOM	985	H	SER	214	-7.705	2.322		1.00 15.00	Ą
ATCM	956	ΞÀ	SER	214	-7.463	1.456		1.00 69.62	Ä
ATOM	957	23	SER	214	8.727	2.218		1.00 67.82	Ą
ATCM	953	03	SER	214	-9.563		-23.336	1.00 67.64	A
ATOM	959	HG	SER	214	-13.468	2.398	-23.623	1.00 15.00	A

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FIGURE 1Q

ATOM	960	C	SER	214	-6.518	1.587	-25.300	1.33	72.29	Ä
MCTA	961	ε	SER	214	-6.102		-25.686		73.45	Ä
	962		٨نن٨	215	-6.175		-25.899	1.23	73.38	
ATOM					-5.456				2 . 2 =	Ä
ATOM.	963	H	جننج	215			-26.565		-3.00	À
ATOM	964	CA	منته	215	-6.858	-0.915	-25.753	1.00	72.62	A
ATOM	965	CB	ALA	215	-7.199	-1.505	-27.138	1.00	73.0E	Ä
ATOM	966	С	ALA	215	-6.331	-2.148	-24.963	1.00	72.11	Ä
ATOM	967	С	۸ننه	215	-7.620	-3.161	-25.069		72.74	Ä
ATOM	968	N	LYS	216	-5.153		-24.262		70.17	, , , , , , , , , , , , , , , , , , ,
	969	н	LYS	216	-4.747		-24.199		15.00	Ç
ATOM										À
ATOM	970	CA	LYS	216	-4.482		-23.626		67.38	Ä
ATOM	971	СB	LYS	216	-3.458		-22.648		65.3C	۸
ATOM	972	CG	LYS	216	-2.217	-2.107	-23.321	1.00	66.B6	A
ATOM	973	CD	LYS	216	-1.419	-3.149	-24.134	1.00	68.81	Ä
ATOM	974	CE	LYS	216	-0.082	-2.674	-24.740	1.00	67.51	Ä
ATOM	975	NZ	LYS	216	0.483		-25.598		67.80	Ä
ATOM	976		LYS	216	0.620	_	-25.041		15.00	Ä
ATOM	977		LYS	216	-0.168		-26.385		15.00	Ä
	978		LYS	216	1.401					
ATOM							-25.973		15.00	À
ATOM	979	C	LYS	216	-5.321		-22.993		66.99	À
ATOM	980	0	LYS	216	-6.462		-22.575		69.90	A
ATOM	981	N	PRO	217	-4.835		-22.952		65.06	Α
ATOM	982	CD .	PRO	217	-3.525	-6.262	-23.308	1.00	67.91	A
ATOM	983	CA	PRO	217	-5.792	-6.827	-22.626	1.00	62.80	A
ATOM	984	CB	PRO	217	-5.285	-8.004	-23.464	1.00	64.33	A
ATOM	985	CG	PRO	217	-3.755		-23.338	1.00	69.63	A
ATOM	986	C	PRO	217	-5.837		-21.150		59.77	Ä
MOTA	987	ō	PRO	217	-4.747		-20.589		58.81	Ä
ATOM	988	N	CYS	218	-7.115		-20.627	_	55.45	
ATOM	989		CYS	218	-7.B74		-21.233			ý
		H							15.00	À
MOTA	990	CA	CYS	218	-7.433		-19.210		46.55	A
MCTA	991	CB	CYS	218	-8.105		-19.079		44.69	Ą
ATOM	992	SG	CYS	218	-8.855		-17.460		43.11	A
ATOM	993	C	CYS	218	-6.265	-7.994	-18.263	1.00	43.24	A
ATOM	994	0	CYS	218	-5.720		-17.959	1.00	44.68	A
ATOM	995	N	GLY	219	-5.853	-6.820	-17.876	1.00	40.28	A
ATOM	996	H	GLY	219	-6.328	-5.961	-18.059	1.00	15.00	A
ATOM	997	CA	GLY	219	-4.659	-6.828	-17.070	1.00	36.27	A
ATOM	998	C	GLY	219	-5.017		-15.643		33.86	Ä
ATOM	999	ō	GLY	219	-5.906		-15.097		34.90	Ä
MOTA	1000	N	GLN	220	-4.313		-15.023		33.15	Ä
ATOM	1001	н	GLN	220	-3.835		-15.580		15.00	
ATOM	1002	CA	GLN	220	-4.448		-13.578			A
				220		17.			29.92	A
ATOM	1003	CB	GLN		-4.298		-12.936		27.81	A
ATOM	1004	CG	GLN	220	-5.380		-11.883		30.94	A
ATOM	1005	CD	GLN	220			-11.132		36.37	A
ATOM	1006	OE:	GLN	220	-4.215		-10.661	1.00	38.47	A
MCTA	1007	NE2	GLN	220	-6.425	-11.296	-10. 9 77	1.00	37.61	Ä
ATOM	1008	HE21	GLN	22C	-6.295	-12.235	-10.667	1.00	15.00	A
ATOM	1009	HE22	GLN	220	-7.373	-11.036	-11.200		15.00	Ä
MOTA	1010	\subseteq	GLN	220	-3.666		-12.859		27.48	A
ATOM	1011	Ċ	GLN	225	-2.461		-12.999		27.61	Ä
ATOM	1010	Ñ	GLN	221	-4.438		-12.110		25.10	
ATOM	1013	H	SLN	:::	-5.433		-12.143			•
ATOM		Ξλ	32%						15.00	÷
	7		·\ ·\		-3.803		-11.387		22.41	À
ATOM		CE	32%	22:	-4.077	-	-11.949		22.12	Ä
ATOM	1316	23	GLN	441	-3.284		-13.163		32.16	Ä
MCTA	1017	20	GLN.	===	- 3 . 795		-13.405		34.69	Ä
ATOM	1018	CE:	SLN	221	-3.746		-12.558		42.12	Ä
MCTA	1019	NEC	321:	==:	-4.648	-1.507	-14.398	1.00	34.93	À

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FIGURE 1R

ATOM	1029	HE2:	. GLN	221	-4.981	-2.187	-15.040	:	Ä
ATOM	1021	HESS		221	-4.944	-0.551	-14.575	. 1.33 18.33	
									÷
ATOM	1022	0	GLN	221	-4.227	-4.913	-9.948	1.00 19.84	Ä
ATOM	1023	Э	GLN:	221	-5.300	-5.381	-9.611	1.00 19.48	Ä
MCTA	1024	N	SER	222	-3.374	-4.330	-9.123	1.00 18.10	A
ATOM	1025	H	SER	222	-2.442	-4.098	-9.441	1.00 15.00	
ATOM	1026	CA	SER	222	-3.851	-4.120	-7.752	1.00 19.45	
			SER	222	-3.104				÷
ATOM	1027	CB				-4.947	-6.691	1.00 19.99	Ä
ATOM	1028	ဝင	SER	222	-3.096	-€.339	-7.053	1.00 24.64	2
ATOM	1029	HG	SER	222	-2.651	-6.336	-7.904	1.00 15.00	Ä
ATOM	1030	C	SER	222	-3.731	-2.688	-7.330	1.00 24.09	÷
ATOM	1031	0	SER	222	-2.992	-1.929	-7.944	1 00 29.41	À
ATOM	1032	N	ILE	223	-4.534	-2.386	-6.283	1.00 22.81	٠ ٨
ATOM	1033	н	ILE	223	-5.172	-3.127	-6.074	1.00 15.00	Ä
		CA.	ILE	223	-4.567				
ATOM	1034					-1.122	-5.530	1.00 21.06	À
MOTA	1035	CB	ILE	223	-5.970	-0.490	-5.852	1.00 19.67	Ä
MOTA	1036		ILE	223	-6.564	0.315	-4.673	1.00 16.59	Ä
ATOM	1037	CG1	ILE	223	-5.911	0.278	-7.188	1.00 15.22	A
ATOM	1038	CD1	ILE	223	-7.229	0.868	-7.709	1.00 20.54	A
ATOM	1039	C	ILE	223	-4.367	-1.446	-4.007	1.00 21.62	A
ATOM	1040	ŏ	ILE	223	-5.098	-2.269	-3.444	1.00 19.58	Ä
		N	HIS	224					
ATOM	1041				-3.429	-0.767	-3.340	1.00 19.73	A
ATOM	1042	H	HIS	224	-2.794	-0.230	-3.899	1.00 15.00	, A
ATOM	1043	CA	HIS	224	-3.497	-0.671	-1.858	1.00 16.45	A
ATOM	1044	CB	HIS	224	-2.164	-1.183	-1.227	1.00 18.74	Α
MOTA	1045	CG	HIS	224	-2.182	-1.442	0.296	1.00 14.92	A
MOTA	1046	ND1	HIS	224	-2.479	-2.628	0.582	1.00 15.33	A
ATOM	1047		HIS	224	-2.667	-3.515	0.505	1.00 15.00	À
ATOM	1048		HIS	224	-1.964				
						-0.524	1.310	1.00 13.79	À
ATOM	1049		HIS	224	-2.137	-1.127	2.517	1.00 10.52	A
MCTA	1050		HIS	224	-2.458	-2.411	2.232	1.00 11.70	À
MOTA	1051	C	HIS	224	-3.914	0.699	-1.284	1.00 15.18	A
ATOM	1052	0	HIS	224	-3.338	1.732	-1.520	1.00 14.36	A
ATOM	1053	N	LEU	225	-4.970	0.673	-0.468	1.00 16.85	A
MOTA	1054	н	LEU	225	-5.317	-0.238	-0.252	1.00 15.00	À
ATOM	1055	CA	LEU	225	-5.395	1.885	0.256	1.00 15.55	Ä
ATOM	1056	CB	LEU	225	-6.927	2.082			
ATOM							0.208	1.00 17.15	À
	1057	CG	LEU	225	-7.495	2.456	-1.154	1.00 18.03	À
MOTA	1058		LEU	225	-6.792	3.659	-1.774	1.00 19.34	A
MOTA	1059		LEU	225	-8.994	2.659	-1.098	1.00 13.66	Ä
ATOM	1060	C	LEU	225	-5.074	1.758	1.739	1.00 14.77	A
ATOM	1061	0	LEU	225	-5.347	0.726	2.345	1.00 12.20	A
ATOM	1062	N	GLY	226	-4.544	2.829	2.344	1.00 18.04	Ä
ATOM	1063	н	GLY	226	-4.218	3.616	1.813	1.00 15.00	_
ATOM	1064	CA	GLY	226	-4.541	2.833			A
							3.841	1.00 18.37	A
ATOM	1065	Č	GLY	226	-4.193	4.171	4.544	1.00 17.08	A
MCTA	1066	0	GLY		-3.389	4.906			Ä
ATOM	1067	N	GLY	227	-4.781	4.457	5.725	1.00 16.30	A
ATOM	1068	н	GLY	227	-5.434	3.771	6.036	1.00 15.00	A
MOTA	1069	CA	GLY	227	-4.379	5.649	6.490	1.00 8.52	A
ATOM	1070	C	GLY	227	-4.935	5.631	7.959	1.00 12.75	Ä
ATOM	1071	č	GLY	227	-5.651	4.748		1.00 10.57	
MCTA	1072	N							A
			VAL	22B	-4.588	6.698	.B.675	1.00 9.23	A
ATOM	1073	H	VAL	228	-4.040	7.398	8.222	1.00 15.00	A
ATOM	1074	CA	VAL	229	-5.110	6.818	10.067	1.00 11.74	Ä
ATOM	1075	CE	VAL	228	-4.085	7.320	11.144	1.00 14.30	A
MOTA	1276	23:	VAL	228	-2.830	6.445		1.00 10.73	*
MOTA	1077	CGS	VAL	228	-4.789	7.565	12.479	1.00 17.07	* * * *
ATOM	:278	2	VAL	228	-6.238	7.803	10.098	1.00 9.03	Â
MOTA	279	Š	VAL	228	-6.089			1.00 12.01	
									,-a

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FIGURE 1S

MCTA	1080		PHE	229	-7.347	7.299	10.640			
ATOM	1081	ä	PHE	229	-7.329	6.332				÷
		~ .					10.922	1.00	15.11	Ä
ATOM	1082	22	PHE	229	- 5 . 56 6	5.106	10.772	1.55	11.15	À
MCTA	1983	CB	PHE	229	- 5 . 578	7.687	9.686	1.33	÷.::	Ä
ATOM	1084	CG	PHE	229	-9.063	7.912	5.233	1.00	5.40	Ä
ATOM	1085	CD:	PHE	229	-9.140	9.196	7.649	1.00	10.53	Ä
MOTA	1086	CDZ		229	-9.433	6.883	7.517			
ATOM	1067	CEI		229	-8.512				6.57	÷
	_					9.443	6.395	1.00	5.18	
ATOM	1098	CE2		229	-7.771	7.128	6.282	1.00	4.26	Ä
MCTA	1089	CZ	PHE	229	-7.813	8.424	5.731	1.00	5	Ä
ATOM	1090	C	PHE	229	-9.202	8.014	12.197	1.00	14.39	A
MOTA	1091	0	PHE	229	-9.116	7.000	12.870	1.00		Ä
ATOM	1092	N	GLU	230	-9.863	9.064	12.672	1.00		Ä
ATOM	1093	Н	GLU	230	-9.912	9.892	12.113	1.00		Ä
ATOM	1094	CA	GLU	230	-10.856	8.944				
	_						13.770	1.00		÷
MOTA	1095	CB	GLU	230	-11.218	10.303	14.393	1.00		À
MOTA	1096	CG	GLU	230	-11.068	10.090	15.889	1.00		Ä
MOTA	1097	CD	GLU	230	-12.314	10.091	16.805	1.00	33.06	Ä
ATOM	1098	OE1	GLU	230	-13.355	10.707	16.552	1.00	38.26	A
ATOM	1099	OE2	GLU	230	-12.218	9.477	17.863	1.00		A
ATOM	1100	С	GLU	230	-12.225	8.268	13.453	1.00		Ä
ATOM	1101	ō	GLU	230	-12.967	€.519	12.492			
ATOM	1102	·N	LEU	231	-12.542			1.00		Á
						7.334	14.361	1.00		A
ATOM	1103	H	LEU	231	-11.840	7.125	15.015	1.00		Α
ATOM	1104	CA	LEU	231	-13.885	6.836	14.330	1.00	13.52	A
MCTA	1105	CB	LEU	231	-13.954	5. 37 8	14.002	1.00	13.90	A
ATOM	1103	CG	LEU	231	-13.199	5.064	12.725	1.00	15.44	A
ATOM	1107	CDI	LEU	231	-13.781	5.712	11.436	⊥.00 :	10.24	A
ATOM	1108		LEU	231	-12.970	3.569	12.769	1.00		Ä
ATOM	1109	C	LEU	231	-14.638	7.074	15.591	1.00		Ä
MCTA	1110	ō	LEU	231	-14.145	6.912	16.692	1.00		
ATOM	1111	N	GLN	232	-15.891	7.411				Ä
ATOM	1112	H	GĽN:	232	-16.107		15.350	1.00		À
	1113					7.560	14.394	1.00		A
ATOM		CA	GLN	232	-16.920	7.509	16.389	1.00 2		A
ATOM	1114	CB	GLK	232	-18.132	8.234	15.804	1.00 2		A
ATOM	1115	CG	GLN	232	-17.792	9.709	15.687	1.00 2	28.60	A
MOTA	1116	CD	GLN	232	-17.625	10.200	17.102	1.00	33.66	A
MCTA	1117	OEl	GLN	232	-18.623	10.472	17.742	1.00	38.08	A
ATOM	1118		g <u>l</u> n	232	-16.380	10.254	17.596	1.00	33.41	A
ATOM	1119	HE21	GLN	232	-15.596	10.186	16.972	1.00	15.00	A
ATOM	1120	HE22	SLN	232	-16.387	10.470	18.576	1.00		A
ATOM	1121	C	GLN	232	-17.402	6.148	16.851	1.00		Ä
ATOM	1122	Ċ	GLN	232	-17.368	5.218	16.052	1.00		Ä
ATOM	1123	N	PRO	233	-17.906	6.013	18.115	1.00		
ATOM	1124	ËD	PRC	233	17.962	7.033				A
ATOM	1125	CA	PRO				19.168	1.00 2		A
				233	-18.570	4.747	18.442	1.00	21.21	Ä
MOTA	1126	CB	PRC	233	-19.013	4.987	19.866	1.00 2		A
MOTA	1127	23	PRO	233	-19.661	6.404	20.339	1.00	20.95	A
ATOM	1128	Ξ.	PRO	233	-19.667	4.417	17.434	1.00	23.66	Α
ATOM	1129	S	PRO	233	-20.275	5.319	16.875	1.00	26.89	A
MOTA	1130	N	SLY	234	-19.731	3.140	17.059	1.00		A
ATOM	1131	H	3 <u>1</u> .	234	-19.082	2.466	17.417		15.00	Ä
MCTA	1132	CA	SLY	234	-23.766	2.767	16.072	1.00		Â
MCTA	1111	Ξ	3 <u>.</u> Y	234	-20.545	3.241	14.625	1.30		•
ATOM		:	<u> </u>	234	-31.299	2.980	13.715			?
ATOM	1134	::		235	-19.405			1.00		A
ATOM		 H	X-X	.235		3.926	14.368	1.00		Ä
					-19.096	4.485	15.135	1.00	15.00	* * * * * *
MOTA	1137	CA	<u>ئــٰ</u> ۂ	235	-18.431	3.515	13.296	1.00		
ATOM	::39	ŢB.	***	235	-18.193	2.042	13.039	1.00	6.55	Ä
ATOM	1139	Ξ	۸ ۱ ۸	235	-18.540	4.160	11.993	1.60	21.96	Ä

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FIGURE 1T

ATOM	1145	2	مُـمَ	235	-18.486	5.385		· · · · · · · · · · · · · · · · · · ·	
							****	1000 26.40	Ä.
ATCM	1141	::	SER	236	-18.699	3.498	10.787	1.00 00 84	Ä
ATOM	1142	H	SER	236	-18.324	4.326	12.254		A.
					-18.630				
MOTA	1143	ΞÀ	SER	23 ć	-15.530	2.227	9.961	1.00 17.60	A
MOTA	1144	ÇB	SER	236	-19.905	1.576	9.160	1.00 14.98	÷
ATOM	1145	CS	SER	236	-20.662	0.908	9.533	1.00 21.35	. ,
MOTA	1146	ΗG	·SER	236	-21.599	0.910	9.647	1.00 15.00	Ä
									^
MCTA	1147	C	SER	236	-17.794	2.538	9.714	1.00 13.65	Ä
ATOM	1148	0	SER	236	-17.939	3.614	8.131	1.00 16.29	À
			VAL	237	-16.986		0.201		
ATOM	1149	N				1.567	8.286	1.00 14.95	A
ATOM	1150	H	VAL	237	-16.764	0.823	9.949	1.00 15.00	Ä
ATOM	1151	CA	VAL	237	-16.201	1.802	7.077		
					· · · · · · · · · · · · · · · · · · ·			1.00 11.41	A
ATOM	1152	CB	VAL	237	-14.681	2.004	7.284	1.00 12.49	Ä
ATOM	1153	CG1	VAL	237	-14.113	0.726	7.939	1.00 13.10	
									À
MOTA	1154	CG2	VAL	237	-14.254	3.396	7.846	1.00 10.27	÷
ATOM	1155	C	VAL	237	-16.468	0.746	6.035	1.00 8.76	÷
ATOM	1156	0	VAL	237	-16.827	-0.363	6.341	1.00 12.84	Ä
ATOM	1157	N	PHE	238	-16.354	1.158	4.773	1.00 12.45	Α
					16.331		_		
ATOM	1158	Н	PHE	238	-16.139	2.128	4.652	1.00 15.00	À
MCTA	1159	CA	PHE	238	-16.521	0.213	3.653	1.00 11.21	A
ATOM	1160	CB	PHE	238	-18.013	0.137			
							3.322	1.00 13.00	Α
ATOM	1161	CG	PHE	238	-18.634	1.468	2.899	1.00 12.17	Α
ATOM	1162	ים	PHE	238	-18.763	1.812	1.518	1.00 12.94	A
ATOM	1163	CD2	PHE	238	-19.135	2.332	3.887	1.00 10.55	Α
MCTA	1164	CEI	PHE	238	-19.407	3.010	1.092	1.00 14.01	A
ATOM	1165	ここ2	PHE	238	-19.786	3.504	3.470	1.00 12.74	A
ATOM	1166	CZ	PHE	238	-19.917	3.836	2.100	1.00 13.17	A
MCTA									
	1167	ε	PHE	238	-15.725	0.582	2.379	1.00 11.20	Ä
ATOM	1168	C	PHE	238	-15.137	1.638	2.267	1.00 B.73	Α
MOTA	1169	N	VAL	239	-15.726	-0.300	1.383	1.00 14.34	
									A
ATOM	1170	H	VAL	239	-16.187	-1.170	1.523	1.00 15.00	A
ATOM	1171	CA	VAL	239	-14.982	0.027	0.154	1.00 14.65	A
MOTA	1172	CB	VAL	239	-13.900	-1.043	-0.162	1.00 14.09	À
ATOM	1173	CG1	VAL	239	-13.004	-1.318	1.038	1.00 14.55	À
MOTA	1174		VAL	239	-13.064				
						-0.594	-1.361	1.00 14.74	À
ATOM .	1175	C	VA:	239	-15.930	0.081	-1.043	1.00 18.32	A
ATOM	1176	0	VAL	239	-16.558	-0.903	-1.369		
								1.00 18.99	A
MOTA	2177	N	ASN	240	-16.000	1.207	-1.707	1.00 19.26	A
ATOM	1178	Н	ASN	240	-15.420	1.947	-1.383	1.00 15.00	A
	1179	CA							
ATOM			ASN	240	-16.613	1.355	-3.031	1.00 21.66	A
MCTA	1180	CB	ASN	240	-16.850	2.856	-3.095	1.00 24.58	À
ATOM	1181	CG	ASN	240	-18.167	3.077	-3.708	1.00 29.09	A
ATOM	1182	051	ASN	24 C	-18.948	2.123	-3.740	1.00 35.44	A
ATOM	1183	ND2	ASN	240	-18.293	4.331	-4.166	1.00 34.71	A
MCTA		HD21			-19.149				
				24C		4.489	-4.657	1.00 15.00	Ä
MCTA	1165	C	ASN	240	-15.6 6 9	0.950	-4.184	1.00 20.96	À
MOTA	1186	0	ASN	24 C	-14.473	1.128	-4.058	1.00 20.99	_
							-4.030		A
MCTA	1187	N	VAL	241	-16.189	0.383	-5.275	1.00 21.52	Α
MOTA	1188	H	VAL	241	-17.182	0.230	-5.295	1.00 15.00	A
		ËA			-15.387				
MCTA	1189		VAL	241		0.439	-6.516	1.00 20.56	Ä
ATOM	1190	CB	VAL	241	-14.581	-0.850	-6.849	1.00 18.02	A
ATOM	1191	CG:	VAL	241	-15.501	-2.058	-7.063		
	>-							1.00 15.06	A
ATOM	1190		VAL	241	-13.597	-1.259	-5.764	1.00 20.05	Ä
MOTA	1193	5	VAL	241	-16.253	5.758	-7.741	1.00 18.88	Ä
	1194	-						1.00 10.00	^
ATOM	54	Ξ		141	-17.441	0.500	-7.819	1.00 18.63	÷
MCTA	1195	:;	THE	242	-15.541	1.162	-8.762	1.00 21.24	Ä
ATOM	1196	H	THE						
A . U.T.				242	-14.704	1.653	-8.486	1.00 15.00	Ä
ATCM	1197	CΆ	THR	242	-15.246	1.476	-10.031	1.00 20.63	A
MCTA	1198	75	THR	242	-15.342	2.269	-10.981	1.00 15.80	
								1.00 15.80	Ä
ATOM	1199	23:	THE	242	-14.035	1.663	-10.953	1.00 17.72	Ä

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FIGURE 1U

ATOM	::::	H31	THE	241	-13.721	1.969	-11.812	1.11 18 11	÷
ATOM	1201	232	THR	242	-18.238	3.730	-11.651		Ä
ATOM	1202	0	THR	242	-16.755	2.240		1.00 18.91	Â
ATOM	1203	Š	THE	242	-17.846	5.196			Â
ATOM	1264	N	ASF	243	-15.923	-0.806		1.00 21.98	
	1205	ä	ASP		-15.057				Ä
ATOM				243			-10.221	1.00 15.00	Ä
ATOM	1206	CÀ	ASP	243	-16.092		-11.628	1.00 21.25	4 4
ATOM	1207	CB	ASP	243	-14.905		-12.594	1.00 02.05	Ä
ATOM	1205	CG	ASP	243	-14.932	-0.954		1.00 25.23	Ä
ATOM	1209	OD1	ASP	243	-14.314	0.051	-13.115	1.00 28.43	Ä
ATOM	1210		ASP	243	-15.588	-1.033	-14.535	1.00 33.00	Ä
ATOM	1211	Ç	ASP	243	-16.123	-3.308	-10.923	1.00 00.35	÷
ATOM	1212	0	ASP	243	-15.148	-4.072	-10.967	1.00 20.43	Ä
ATOM	1213	N	PRO	244	-17.204		-10.154	1.00 19.52	Ä
ATOM	1214	CD	PRO	244	-18.481		-10.071	1.00 16.83	Ä
ATOM	1215	CA	PRO	244	-17.120	-4.706	-9.269	1.00 19.13	
ATOM	1216	CB	PRO	244	-18.293	-4.535			÷
ATOM	1217	CG	PRO					1.00 15.33	À
	1218			244	-18.890	-3.174	-8.634	1.00 15.21	À
ATOM		C	PRO	244	-16.975	-6.034	-9.974	1.00 19.29	Ä
ATOM	1219	0	PRO	244	-16.194	-6.859	-9.548	1.00 23.48	A
ATOM	1220	N	SER	245	-17.581		-11.150	1.00 22.60	A
ATOM	1221	Н	SER	245	-18.220		-11.473	1.00 15.00	Ä
MOTA	1222	CA	SER	245.	-17.414		-11.942	1.00 25.50	A
MOTA	1223	CB	SER	245	-18.256	-7.369	-13.234	1.00 21.36	A
ATOM	1224	OG	SER	245	-19.667	-7.567	-12.981	1.00 38.26	A
MCTA	1225	HG	SER	245	-19.848		-12.038	1.00 15.00	A
ATOM	1226	С	SER	245	-15.955		-12.328	1.00 24.14	Ä
MCTA	1227	C	SER	245	-15.477		-12.623	1.00 24.84	Ä
ATOM	1228	N	GLN	246	-15.177		-12.385	1.00 28.52	Ä
ATOM	1229	н	GLN	246	-15.638		-12.265	1.00 15.00	Ä
MOTA	1230	CA	GLN	246	-13.743		-12.590	1.00 26.45	
MOTA	1231	CB	GLN.	246	-13.144		-13.233		Ä
MCTA	1232	CG	GLN	246	-13.403			1.00 29.90	À
MCTA	1233	CD	GLN				-14.758	1.00 26.84	À
ATOM	1234		GLN	246	-14.862		-15.129	1.00 21.60	À
	1235			246	-15.538		-14.616	1.00 24.20	A
ATOM			GLN	246	-15.334		-15.975	1.00 26.15	Ä
ATOM	1230	HE21	GLN	246	-14.763		-16.423	1.00 15.00	À
ATOM	-23/	HE22		246	-16.320		-16.084	1.00 15.00	A
MOTA	1238	Ξ	GLN	246	-12.936		-11.363	1.00 27.14	A
MCTA	1239	0	GLN	246	-11.721		-11.454	1.00 25.73	A
MCTA	1240	N	VAL	247	-13.615		-10.196	1.00 23.70	Ä
MOTA	1241	н	VAL	247	-14.600	-7.594	-10.146	1.00 15.00	A
MOTA	1242	CA	VAL.	247	-12.718	-7.569	-9.097	1.00 21.91	A
ATOM	1243	CB	VAL	247	-13.156	-6.B14	-7.859	1.00 21.59	A
ATOM	1244	CG1	VAL	247	-14.027	-7.616	-6.962	1.00 24.52	À
ATOM	1245	CG2	VA.	247	-13.680	-5.409	-B.167	1.00 21.61	۸
ATOM	1246	Ξ.	VAL	247	-12.258	-8.998	-8.910	1.00 21.55	Ä
ATOM	1247	С	VAL	247	-12.946	-9.912	9.251	1.00 19.53	Ä
MOTA	1248	N	SER	248	-11.000	-9.152	-8.444	1.00 21.31	Ä
ATOM	1245	H	SER	248	-10.558	-8.342	-B.070	1.00 15.00	Ä
ATOM	1250	CA	SER	248	-10.414	-10.499	-8.327		Ä
ATOM	1251	CB	SER	248				1.00 21.97	
ATOM	1252	25 25				-10.571	-8.828	1.00 23.61	À
ATOM	1253		SER	248	-8.860		-10.128	1.00 20.21	À
ATOM	1254	#G	SER	248	-9.752	-10.027	-10.496	1.00 15.00	À
7.07.		:	SER	148	.10.538	-11.576	-6.946	.1.00 19.28	Ä
ATOM	1155	2	SER	248	-10.046	-10.409	-5.052	1.00 20.64	Ä
ATOM	1135	N	HIS	249		-12.204	-6.814	1.00 18.72	Ä
ATOM	::::	Ξ.	#:5	249	-11.284	-12.753	-7.674	1.00 15.00	Ä
MOTA	:::::	24	HIS	249		-12.673	-5.478	1.00 17.21	÷
ATOM	::::	CE	HIS	249	-13.380	-13.152	-5.484	1.00 13.15	A

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FIGURE 1V

ATOM	1260	23	HIS	249	-13.919	-11.905	-5.550	1.21 11.13	À
ATOM	1261	::::	HIS	249	-14.137	-11.129	-1 486	1311 13.47	A
7.00	1260	#5:	HIS	249	-13.720	-11.294	-3.611	11111111111	
ATOM					-14.662	-11.414			â
ATOM	1263		HIS	249			-6.610		â
ATOM	1264		HIS	249		-10.347	-6.134	1.00 15.51	
ATOM	1255	CEI	HIS	245		-10.142	-4.821	1.00 12.36	À
MOTA	1266	C	HIS	249		-13.683	-4.850	1.00 23.58	÷
ATOM	1267	5	HIS	245		-14.729	-4.359	1.00 21.98	A
ATOM	1268	N	GLY	250	-9.398	-13.258	-4.878	1.00 29.10	Ä
ATOM	1269	H	GLY	250		-12.351	-5.253	1.00 15.00	Ä
ATOM	1270	CA	GLY	25C		-14.041	-4.115	1.00 24.27	Ä
	1271	Ξ.	GLY	250		-15.372	-4.743	1.00 25.93	Ä
MOTA				250		-15.520	-5.795	1.00 29.26	, , , , , , , , , , , , , , , , , , ,
ATOM	1272	C	GLY						
ATOM	1273	N	THR	251		-16.302	-4.127	1.00 22.38	÷
ATOM	1274	H	THR	251		-17.038	-4.804	1.00 15.00	À
ATOM	1275	CA	THR	251		-16.139	-2.725	1.00 21.12	À
ATOM	1276	CB	THR	251	-6.988	-17.525	-1.933	1.00 24.76	Ä
ATOM	1277	OG1	THR	251	-5.877	-17.641	-0.981	1.00 22.90	À
ATOM	1278	HG:	THR	251	-6.063	-18.366	-0.381	1.00 15.00	Ä
ATOM	1279	CG2	THR	251		-18.722	-2.890	1.00 22.77	A
ATOM	1280	C	THR	251		-15.158	-2.473	1.00 17.96	À
ATOM	1281	ŏ	THR	251		-15.043	-3.213	1.00 12.30	Ä
	1282	N N	GLY	252		-14.367	-1.419	1.00 16.85	Ä
ATOM						-14.432			Ä
MOTA	1283	H	GLY	252	·		-0.862	1.00 15.00	
MCTA	1284	CA	GLY	252		-13.375	-0.928	1.00 13.16	À.
ATOM	1285	C	GLY	252		-12.058	-1.670	1.00 15.51	À
ATOM	1286	0	GLY	252		-11.168	-1.439	1.00 15.18	Ä
ATOM	1287	N	PHE	253	-6.189	-12.063	-2.744	1.00 16.66	Ä
MCTA	1298	H	PHE	253	-6.868	-12.805	-2.761	1.00 15.00	A
MCTA	1289	CA	PHE	253	-6.110	-10.892	-3.651	1.00 15.77	A
ATOM	1290	CB	PHE	253		-11.216	-5.100	1.00 17.11	Ä
MCTA	1291	ËĞ	PHE	253		-11.840	-5.994	1.00 11.82	A
ATOM	1292	551	PHE	253		-11.175	-6.231	1.00 13.69	Ä
			PHE	253		-13.089	-6.558	1.00 18.59	À
MOTA	1293	CD2						-	Ä
ATOM	1294	CEI	PHE	253		-11.771	-6.993	1.00 14.39	÷
ATOM	1295	CE2	PHE	253	-4.840	-13.680	-7.363	1.00 21.37	
MOTA	1296	CZ	PHE	253		-13.014	-7.562	1.00 15.72	À
ATOM	1297	C	PHE	253	-6.740	-9.599	-3.147	1.00 13.88	À
ATOM	1298	C	PHE	253	-6.347	-8.477	-3.453	1.00 14.27	Ä
MCTA	1299	N	THR	254	-7.865	-9.837	-2.502	1.00 14.00	À
MCTA	1300	H	THR	254	-e.c7 9	-10.748	-2.124	1.00 15.00	À
MCTA	1301	CA	THR	254	-8.741	-8.681	-2.185	1.00 14.09	A
MCTA	1302	CB	THR	254	-9.908	-8.469	-3.201	1.00 11.66	A
ATOM	1303	0G1	THR	254	-9.414	-8.325	-4.536	1.00 13.08	A
ATOM	1304	HG1	THR	254	-9.926	-9.054	-4.992	1.00 15.00	Ä
MCTA	1305	CG2	THR	254	-10.882	-7.321	-2.885	1.00 13.78	Ä
ATOM	1306	_	THR	254		- B . 779			
	1300	~	THR	254	-9.906	-9.695	-0.240	1 00 14.54	Ä
ATOM	1307	0							Â
ATOM	1308	N	SER	255	-9.307		-0.027		<u>~</u>
ATOM	1309	H	SER	255	-8.425	-7.021	-0.490		÷
ATOM	1310	CA	SER	255	-9.032	-7.725	1.431	1.00 7.59	Ä
ATOM	1311	CB	SER	255	•7.793	-8.466	1.976	1.00 6.39	A
MCTA	1312	CG	SER	255	-6.704	-7.56C	2.041	1.00 9.69	'n
ATOM	1313	НS	SER	255	-5.920	-8.031	1.741	1.00 15.00	Ä
ATOM		-	SER	255	-9.248	-6.341	2.085	1.00 10.05	Ä
ATOM	1314	:	SER		-9.191	-5.254	1.492	1.00 15.21	÷
ATOM	1314	::	PHE	256	-9.653	-6.385	3.369		A
ATOM		Ξ.	PHE	156	- 9.700	-7.323	3.733	1.00 15.00	Ä
	1318		PHE	256	-10.114	-5.168	4.035	1.00 7.94	
ATOM				- 3 5	744.414	-2.100			
ATOM	1319	ΞΞ	PHE	256	-11.605	-5.009	3.679	1.00 11.65	Ä

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FIGURE 1W

	1320	25	PHE	. 156	-12.376	-3.524	4.235	1.:: 8		
MCTA									-:	Ė
ATOM	1321	55:	PHE	156	-11.766	-2.570	4.533			Ξ
ATOM	1322	C D2	PHE	256	-13.756	-3.976	4.327	i.:: 6	. = =	Ä.
ATOM	1323	CE:	PHE	256	-12.503	-1.490	5.034		. 4.5	Ä
	1324	SE2		256	-14.514	-2.849	4.734		. : :	
ATOM						•				÷
MCTA	1325	ΞZ	PHE	256	-13.862	-1.657	5.211	1.00 9		Ä
ATOM	1326	C	PHE	256	- 9 . 933	-5.268	5.560	1.00 11	. 92	Ä
MOTA	1327	၁	PHE	25€	-10.195	-6.290	6.177		. 43	À
	_		GLY	257	-9.420	-4.207	6.169	1.00 10		Ξ
ATOM	132B	N								
ATOM	1329	H	GLY	257	-9.217	-3.365	5.653	1.00 15	. OC	à
MOTA	1330	CA	GLY	257	-9.368	-4.406	7.612	1.00 11	.2€	À
ATOM	1331	2	GLY	257	-8.965	-3.122	8.287	1.00 11		Ä
			GLY	257	-8.916	-2.068	7.679			$\hat{\cdot}$
ATOM	1332	0						1.00 10		÷
ATOM	2333	·N	LEU	258	-8.688	-3.277	9.565	1.00 12		Ä
ATOM	1334	н	LEU	258	-8.776	-4.204	9.943	1.00 15	. OC	Ä
ATOM	1335	CA	LEU	258	-8.434	-2.098	10.426	1.00 14	7.7	À
ATOM	1336	CB	LEU	258	-9.751	-1.212	10.704	1.00 14		À
ATOM	1337	CG	LEU	258	-10.991	-1.863	11.379	1.00 18	.G2	À
MOTA	1338	CD1	LEU	258	-12.317	-1.125	11.094	1.00 15	. 05	Ä
ATOM	1339		LEU	258	-10.743	-2.047	12.905	1.00 15		A
ATOM	1340	C	LEU	258	-7.737	-2.525	11.709	1.00 11		λ
ATOM	1341	0	LEU	258	-7.851	-3.690	12.096	1.00 7	.91	Α
ATOM		N	LEU	259	-7.058	-1.537	12.343	1.00 11	. 64	À
ATOM	1343	Н	LEU	259	-6.883	-0.685	11.844	1.00 15		Α
ATOM	1344	CA	LEU	259	-6.581	-1.780	13.714			÷
ATOM	1345	CB	LEU	259	-5.155	-2.417	13.831	1.00 7	.40	Α
ATOM	1346	CG	LEU	259	-4.194	-1.621	12.931	1.00 11	.40	À
MCTA	:347	CD1	LEU	259	-3.355	-2.412	11.926			À
ATOM	1348		LEU	259	-3.379	-0.670	13.808	1.00 13		À
ATOM	1349	C	LEU	259	-6.652	-0.497	14.531	1.00 10	.40	A
ATOM	1350	0	LEU	259	-6.202	0.556	14.082	1.00 9	. 73	A
MCTA	1351	N	LYS	260	-7.193	-0.629	15.762	1.00 12		À
MOTA	1352	H	LYS	260	-7.395	-1.553	16.115	1.00 15		Å
MCTA	1353	CA	LYS	260	-7.069	0.521	16.693	1.00 13	.51	Α
ATOM	1354	CB	LYS	260	-8.014	0.312	17.885	1.00 13	.49	A
ATOM	1355	CG	LYS	260	-8.378	1.656	18.521	1.00 17		A
	1356	SD	LYS	260	-9.435	1.456	19.596	1.00 12		
ATOM										A
MOTA	1357	CE	LYS	260	-10.151	2.681	20.121	1.00 11		A
MCTA	1358	NZ	LYS	260	-9.175	3.595	20.697	1.00 13	. 3 3	A
MCTA	1359	HZ1	LYS	260	-0.534	3.932	19.954	1.00 15	.00	Α
ATOM	1360	HZ2	LYS	260	-9.693	4.404	21.095	1.00 15		A
MOTA	1361		LY5	260	-8.638	3.136	21.458	1.00 15		A
ATOM	1362	C	LYS	260	-5.648	0.921	17.125	1.00 16	. 54	A
MOTA	13£3	0	LY5	260	-4.828	0.112	17.481	1.00 15	. 61	Α
MCTA	1364	N	LEU	261	-5.353	2.199	17.015	1.00 14		Ä
	1365	ä	LEU	261	-5.389	2.838	16.856			
ATOM								1.00 15		À
ATOM	1355	CB	LEU	251	-3.705	4.005	17.185	1.00 19	. 53	A
ATOM	1367	ca	LEU	261	-3.177	4.309	15.787	1.00 16	.82	Ä
MCTA	1368		LEU	261	-3.010	5.779	15.767	1.00 12		Ä
ATOM	1369		LEU	261	-4.010	3.906	14.577	1.00 18		Ä
MOTA	1370	2	LEU	261	-4.243	2.667	19.225	1.00 20	.80	Ä
ATOM	1371	CCT: CCT:	LEU	261	-5.363	2.741	19.746	1.00 22	.59	À
ATOM	1372	2277	LEU	261	-3.221	2.696	19.913	1.00 26		Ä
ATOM	13-3	CA	LEU	261	-4.122	2.604	17.684	1.00 18		~
		-^								À
ATOM	1374	:	HCH	531	-20.040	9.837	7.596	1.00 16		~
ATOM	1375	H1	HCH	501	-19.411	10.547	7.803	1.00 10	. ၁၀	×
ATOM	:374	H.C	HCH	50:	-19.615	9.317	6.900	1.00 10		×
ATOM	13	5	HIH	5:2	-9.727	11.545	10.743	1.00 10		×
		<u>.</u>		=						
ATOM	1378	#:	HCH	502	-10.039	11.934	9.919	1.00 15		~
ATOM	1379	HI.	HCH	500	-11.233	12.125	11.315	1.00 15	.00	₩

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FIGURE 1X

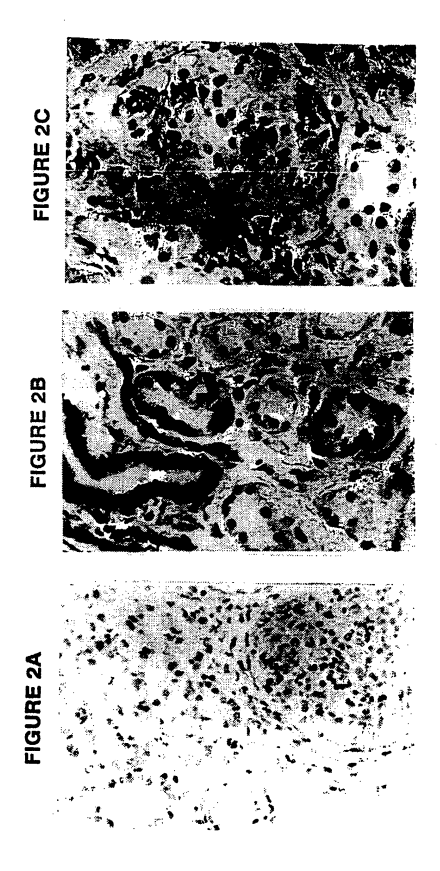
ATOM	1380	S	HOH	503	-5.158	13.168	13.681		
								1.00 30.64	₩.
MOTA	1381	#1	HOH	503	-8.715		13.277	1.00 15.00	₩.
MCTA	1362	H2	HOH	503	-8.700	13.944	13.574	1.00 15.00	¥
ATOM	1383	C	HOH	504	-16.772	8.440	12.789	1.00 12.00	×
MCTA	1384	H1	нон	504	-17.194				
							12.886	1.00 10.00	¥
ATOM	1385	H2	HOH	504	-15.921		12.582	1.00 10.00	×
ATOM	1386	0	нон	505	-25.173	7.297	7.925	1.00 47.03	₩
ATOM	1387	H1	нон	505	-24.690		B.239	1.00 10.00	
								1.00 10.00	¥
ATOM	1388	H2	HOH	505	-25.990		7.583	1.00 10.00	×
ATOM	1389	0	нон	506	-23.612	14.948	13.859	1.00 36.14	₩
ATOM	1390	H1	HOH	506	-24.160	15.702	13.605	1.00 10.00	×
	1391	H2	нон	506	-23.282				
ATOM							14.748	1.00 10.00	W
ATOM	1392	0	HOH	507	-17.329		-7.186	1.00 34.02	₩.
ATOM	1393	C	нон	508	-18.687	-7.253	-3.843	1.00 63.14	W
ATOM	1394	0	HOH	509	-7.157	11.327	3.239	1.00 22.26	¥
ATOM	1395	ō	НОН	510	-19.322				
		_					-2.227	1.00 37.69	₩
ATOM	1396	0	HOH	511	-14.645	-7.711	-1.931	1.00 26.48	W
ATOM	1397	0	нон	512	-18.377	-9.754	12.556	1.00 24.86	W
ATOM	1398	0	HOH	513	0.030	O OAR	-13.455	1.00 26.05	W
MOTA	1399	0	нон	514	-8.938		22.862	1.00 34.39	W
MOTA	1400	0	HOH	515	-29.446	-4.922	-7.247	1.00 41.61	W
ATOM	1401	0	HOH	516	-12.982	10.220	10.038	1.00 47.16	W
MOTA	1402	0	HOH	517	-21.797		7.242	1.00 60.65	
									W
MOTA	1403	0	нон	518	-7.867	8.165	19.484	1.00 40.46	₩
ATOM	1404	၁	HOH	520	-15.588	-14.701	14.628	1.00 63.80	W
ATOM	1405	0	нон	521	-21.844	7.778	20.415	1.00 35.72	W
ATOM	1406	0	HOH	522	-6.555	-3 308	-15.790	1.00 33.63	
									W
ATOM	1407	0	HOH	523		-13.476	-8.051	1.00 44.08	W
ATOM	1408	၁	нон	524	-17.413	-9.311	17.071	1.00 34.06	W
ATOM	1409	0	HOH	525	-23.838	4.781	19.884	1.00 37.99	W
ATOM	1410	C	нон	526	-26.323	15.525	10.379	1.00 72.49	W
MCTA	1411	0	нон	527		-13.749		1.00 43.99	W
ATOM	1412	С	HOH	528	-0.470	2.513	17.943	1.00 63.68	W
ATOM	1413	0	нон	529	-5.580	-12.778	-14.864	1.00 47.52	W
MCTA	1414	0	HOH	530	-2.641	7.004	2.495	1.00 18.07	W
ATOM	1415	ō	нон	531	-6.472				
								1.00 24.96	W
MOTA	1416	0	HOH	532		-16.426	-0.360	1.00 63.56	W
ATOM	1417	0	HOH	533	-1.378	-17.183	-13.053	1.00 67.67	W
MOTA	1418	0	HOH	534	-4.774	9.073	-0.651	1.00 23.36	w
MOTA	1419	ō	нон	535		-13.857	6.913		
								1.00 32.28	W
ATOM	1420	C	нон	536	-23.062	3.270	0.454	1.00 52.03	W
ATOM	1421	0	HOH	537	-25.906	9.022	16.986	1.00 44.75	W
ATOM	1422	0	HOH	538	-21.729	16.972	17.027	1.00 53.12	W
ATOM	1423	0	HOH	539	-9.0B4	11.806	17.034	1.00 70.90	W
		0							
MCTA	1424		нон	54 C		-13.296	15.207	1.00 35.65	W
ATOM	1425	C	нон	541	-6.968	13.255	17.989	1.00 67.36	W
ATOM	1426	0	HOH	542	-20.593	-11.039	-9.003	1.00 96.30	W
ATOM	1427	0	HOH	543	-15.926		1.269	1.00 35.72	W
ATOM	1428								
		C	нон	544	-24.591	-7.285	-2.353	1.00 43.42	W
ATOM	1429	၁	нон	545	- 25 . 859	-2.666	-15.747	1.00 53.56	W
ATOM	1430	0	HOH	546	-23.074	-1.533	11.026	1.00 56.44	W
ATOM	1431	0	HOH	548	-8.941		-12.394	1.00 64.34	W
	1432	~							
MOTA		_	нон	549	-14.150		-12.250	1.00 41.38	W
MCTA	1423	Ξ	нон	550	-14.274	-0.613	18.441	1.00 56.17	w
ATOM	1434	Ĵ	HOH	551	-12.241	-19.609	8.637	1.00 80.90	W
ATOM	1435	Ξ	HOH	552	-10.316	15.578	10.166	1.00 39.58	W
MCTA	1436	_	нон	553					
		000000			-15.367	10.941	14.659	1.00 40.40	W
ATOM	1437	Ξ	нон	554	-2.322	1.830	-5.294	1.00 33.65	w
ATOM	1438	0	нон	555	-22.393	-14.875	-4.217	1.00 52.40	w
ATOM	1439	2	HOH	556	-22.120	14.279	7.189	1.00 38.55	W
		-						2.00 30.33	71

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FIGURE 1Y

MCTA	1440	C	HOH	557	-28.533	6.135	9.560	1.00 37.40	~
ATOM	1441	S	нон	556	-5.554	-16.509	13.192	1.11 88.88	*
ATOM	1442	Ĉ	HOH	559	-22.996	12.522	1.162	1:00 63 77	₩.
ATOM	1443	ō	нон	560	-13.764	2.268	-14.743	1.00 27.47	∵
ATOM	1444	č	нон	561	-15.556	7.750	-5.628	1.00 75.65	¥
	1445	Ö	HOH	562	-1.970	-15.363	-17.719	1.00 76.30	₩.
ATOM	1446	Ö	нон	563	-18.939	-0.335	-13.842	1.00 48.39	¥
ATOM	1445	0	нон	564	.12.619	14.760	-6.974	1.00100.59	¥
ATOM	_	Ö	нон	565	-9.491	18.046	13.682	1.00 87.45	₩
MOTA	1448		HOH	566	-11.655	-11.140	22.481	1.00 28.88	W
MOTA	1449	0	нон	567	-24.072	-3.264	-0.332	1.00 35.13	W
ATOM	1450	C		568	-27.455	0.119	-7.117	1.00 71.07	×
MOTA	1451	0	нон		-14.604	3.516	-6.119	1.00 59.45	w
MOTA	1452	0	нон	569	-2.635		-16.973	1.00 59.09	×
AŢOM	1453	0	нон	570			-7.543	1.00 34.10	ŵ
ATOM	1454	0	HOH	571	-18.841	4.066	17.953	1.00 70.45	×
MOTA	1455	0	нон	572	-24.996	1.301		1.00 70.45	W.
ATOM	1456	0	HOH	573	-14.666	16.471	8.995		W
ATOM	1457	0	HOH	574	-14.786	1.426	10.949	1.00 82.68	¥
ATOM	1458	0	HOH	575		-14.717	-4.352	1.00 29.09	W
ATOM	1459	0	HOH	576	-16.273		6.109	1.00104.64	W
ATOM	1460	0	HOH	577	-25.471	-0.127	-2.510	1.00 62.74	W
ATOM	1461	0	HOH	578		-17.173	19.514	1.00 89.62	
ATOM	1462	0	HOH	579	-21.060	14.259		1.00 69.59	W
MOTA	1463	0	HOH	580	-19.286		-12.816	1.00 60.37	W
ATOM	1464	0	HOH	581		-15.840	0.317	1.00 58.24	W
ATOM	1465	0	HOH	582	-22.434	-10.539	12.489	1.00 70.25	W
ATOM	1466	0	нон	583	-21.327	3.668	-2.500	1.00 39.32	W
ATOM	1467	С	HOH	584	-25.325	5.247	16.919	1.00 41.31	W
ATOM	1468	0	HOH	585	-24.945	-10.718	-2.375	1.00 38.85	W
ATOM	1469	С	HOH	586	-24.342	-13.003	1.927	1.00 70.58	W
MCTA	1470	C	НОН	587	-18.020	11.871	11.358	1.00 64.47	₩
ATOM	1471	Ó	нон	588	-27.135	6.965	13.151	1.00 53.96	W
ATOM	1472	0	нон	589	-14.982	-16.230	-2.494	1.00 30.24	W
ATOM	1473	Ö	нон	590	-5.646	14.418	-2.232	1.00 41.78	W
ATOM	1474	Č	нон	591	-2.745	-0.153	-17.104	1.00 55.19	W
ATOM	1475	č	нон	592	-3.397	-7.012	22.477	1.00 59.46	w
ATOM	1476	ŏ	нон	593	-32.916	-4.705	-4.143	1.00 51.88	W
ATOM	1477	5	нон	594	-10.913	-18.855	-3.503	1.00 42.29	₩
ATOM	1478	Õ	нон	595	-24.157	1.821	-6.165	1.00 47.43	W
END	44.0	•			- · - -				
ط11ت									



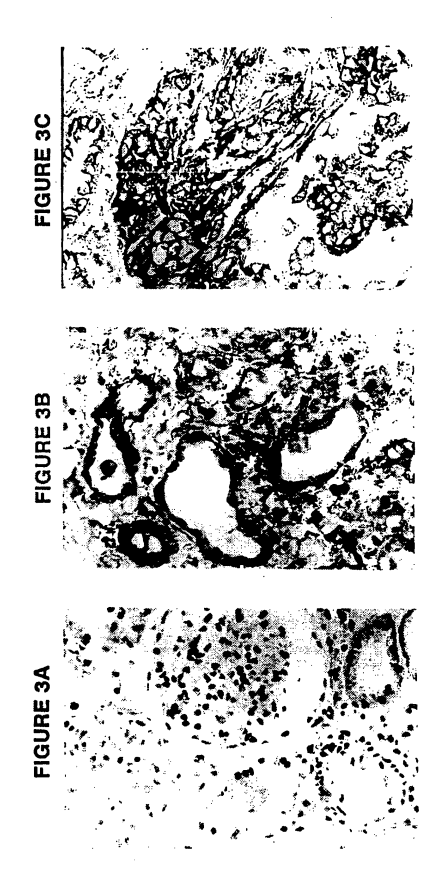






FIGURE 4B



FIGURE 5

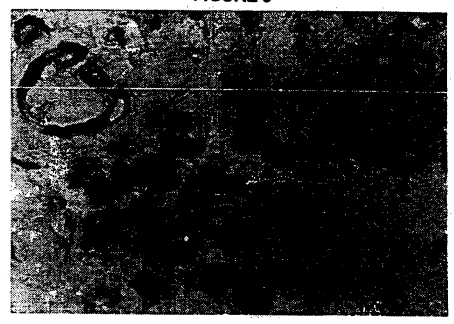


FIGURE 6

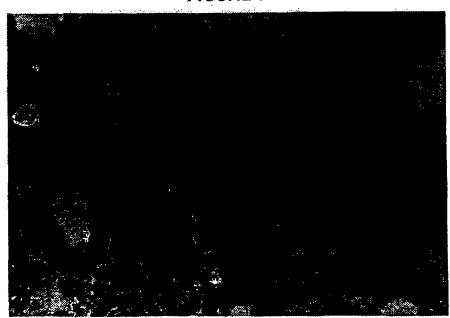
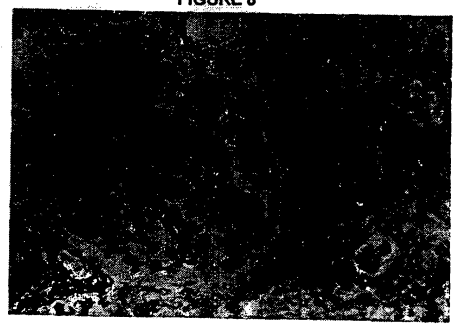






FIGURE 8







INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/00668

A. CLASSIFICATION OF SUBJECT MATTER	
IPC(6) :A61K 38/02, 38/17; 39/395 US CL :424/130.1, 133.1, 141.1, 144.1, , 154.1, 173.1; 514/2, 8	
According to International Patent Classification (IPC) or to both national classification and IPC	
B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols)	
· · · · · · · · · · · · · · · · · · ·	
U.S. : 424/130.1, 133.1, 141.1, 144.1, . 154.1, 173.1; 514/2, 8	
Documentation searched other than minimum documentation to the extent that such documents are included	I in the fields searched
Electronic data base consulted during the international search (name of data base and, where practicable	, search terms used)
APS, DIALOG, BIOSIS, CA, EMBASE, MEDLINE, WPI search terms: 5c8, cd40L, cd40 ligand, kidney, renal	
C. DOCUMENTS CONSIDERED TO BE RELEVANT	
Category* Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
WO 93/09812 A1 (THE TRUSTEES OF COLUMBIA UNIESTLY IN THE CITY OF NEW YORK) 27 MAY 1993, see entire document.	1-23, 31-57, 65-76
	24-30, 58-64
Kidney International, Volume 48, issued 1995, Biancone et	1-23, 31-57,
al., "Inhibition of the CD40-CD40 ligand pathway prevents	65-76
murine membranous glomerulonephritis", pages 458-468,	
see entire document.	24-30, 58-64
Structure, Volume 3, issued 15 October 1995, Karpusas et al., "2 A crystal structure of an extracellular fragment of human CD40 ligand", pages 1031-1039, see entire document.	1-76
X Further documents are listed in the continuation of Box C. See patent family annex.	
Special categories of cited documents: "T" later document published after the inter-	mational filing date or priority
document defining the general state of the art which is not considered to be of particular relevance date and not in conflict with the application to be of particular relevance	stion but cited to understand the
cartier document published on or after the international filling date "X" document of particular relevance; the considered novel or cannot be considered novel or cannot be considered when the document is taken alone	e claimed invention cannot be red to involve an inventive step
cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the considered to involve an inventive combined with one or more other suc-	step when the document is documents, such combination
document published prior to the international filing date but later than "&" document member of the same patent the priority date claimed	ne art
ate of the actual completion of the international search Date of mailing of the international sea	rch report
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01 MAY 1997 Ø2 JUN 1997	· -

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US97/00668

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT								
Category*	ategory* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim N							
Y	Science, Volume 257, issued 21 August 1992, Kuntz e "Structure-Based Strategies for Drug Design and Disco pages 1078-1082, see entire document.	et al., overy",	1-76					
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